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06 January 2017

Version of attached file:

Published Version

Peer-review status of attached file:

Peer-reviewed

Citation for published item:

Mason, A. R. and Mason, J. and Cork, M. and Dooley, G. and Hancock, H. (2013) 'Topical treatments for chronic plaque psoriasis.', Cochrane database of systematic reviews. (3).

Further information on publisher's website:

https://doi.org/10.1002/14651858.CD005028.pub3

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Topical treatments for chronic plaque psoriasis (Review)

Mason AR, Mason J, Cork M, Dooley G, Hancock H

Mason AR, Mason J, Cork M, Dooley G, Hancock H. Topical treatments for chronic plaque psoriasis. *Cochrane Database of Systematic Reviews* 2013, Issue 3. Art. No.: CD005028. DOI: 10.1002/14651858.CD005028.pub3.

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[Intervention Review]

Topical treatments for chronic plaque psoriasis

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Editorial group: Cochrane Skin Group. **Publication status and date:** Edited (no change to conclusions), published in Issue 4, 2013. **Review content assessed as up-to-date:** 2 February 2011.

Citation: Mason AR, Mason J, Cork M, Dooley G, Hancock H. Topical treatments for chronic plaque psoriasis. *Cochrane Database of Systematic Reviews* 2013, Issue 3. Art. No.: CD005028. DOI: 10.1002/14651858.CD005028.pub3.

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ABSTRACT

Background

Chronic plaque psoriasis is the most common type of psoriasis, and it is characterised by redness, thickness, and scaling. First-line management of chronic plaque psoriasis is with topical treatments, including vitamin D analogues, topical corticosteroids, tar-based preparations, dithranol, salicylic acid, and topical retinoids.

Objectives

To compare the effectiveness, tolerability, and safety of topical treatments for chronic plaque psoriasis, relative to placebo, and to similarly compare vitamin D analogues (used alone or in combination) with other topical treatments.

Search methods

We updated our searches of the following databases to February 2011: the Cochrane Skin Group Specialised Register, CENTRAL in *The Cochrane Library* (2011, Issue 2), MEDLINE (from 1948), EMBASE (from 1980), Science Citation Index (from 2008), Conference Proceedings Citation Index - Science (from 2008), BIOSIS (from 1993), Dissertation Abstracts via DialogClassic (all publication years), and Inside Conferences (all publication years).

We identified ongoing and unpublished studies from the UK Clinical Research Network Study Portfolio and the *meta*Register of Controlled Trials. We checked the bibliographies of published studies and reviews for further references to relevant trials, and we contacted trialists and companies for information about newly published studies.

A separate search for adverse effects was undertaken in February 2011 using MEDLINE and EMBASE (from 2005).

Final update searches for both RCTs and adverse effects were undertaken in August 2012. Although it has not been possible to incorporate RCTs and adverse effects studies identified through these final searches within this review, we will incorporate these into the next update.

Selection criteria

Randomised trials comparing active topical treatments against placebo or against vitamin D analogues (used alone or in combination) in people with chronic plaque psoriasis.

Topical treatments for chronic plaque psoriasis (Review)

Data collection and analysis

One author extracted study data and assessed study quality. A second author checked these data. We routinely contacted trialists and companies for missing data. We also extracted data on withdrawals and on local and systemic adverse events. We defined long-term trials as those with a duration of at least 24 weeks.

Main results

This update added 48 trials and provided evidence on 7 new active treatments. In total, the review included 177 randomised controlled trials, with 34,808 participants, including 26 trials of scalp psoriasis and 6 trials of inverse psoriasis, facial psoriasis, or both. The number of included studies counted by Review Manager (RevMan) is higher than these figures (190) because we entered each study reporting a placebo and an active comparison into the 'Characteristics of included studies' table as 2 studies.

When used on the body, most vitamin D analogues were significantly more effective than placebo, with the standardised mean difference (SMD) ranging from -0.67 (95% CI -1.04 to -0.30; 1 study, 119 participants) for twice-daily becocalcidiol to SMD -1.66 (95% CI - 2.66 to -0.67; 1 study, 11 participants) for once-daily paricalcitol. On a 6-point global improvement scale, these effects translate into 0.8 and 1.9 points, respectively. Most corticosteroids also performed better than placebo; potent corticosteroids (SMD -0.89; 95% CI -1.06 to -0.72; I² statistic = 65.1%; 14 studies, 2011 participants) had smaller benefits than very potent corticosteroids (SMD - 1.56; 95% CI -1.87 to -1.26); I² statistic = 81.7%; 10 studies, 1264 participants). On a 6-point improvement scale, these benefits equate to 1.0 and 1.8 points, respectively. Dithranol, combined treatment with vitamin D/corticosteroid, and tazarotene all performed significantly better than placebo.

Head-to-head comparisons of vitamin D for psoriasis of the body against potent or very potent corticosteroids had mixed findings. For both body and scalp psoriasis, combined treatment with vitamin D and corticosteroid performed significantly better than vitamin D alone or corticosteroid alone. Vitamin D generally performed better than coal tar, but findings relative to dithranol were mixed. When applied to psoriasis of the scalp, vitamin D was significantly less effective than both potent corticosteroids and very potent corticosteroids. Indirect evidence from placebo-controlled trials supported these findings.

For both body and scalp psoriasis, potent corticosteroids were less likely than vitamin D to cause local adverse events, such as burning or irritation. Combined treatment with vitamin D/corticosteroid on either the body or the scalp was tolerated as well as potent corticosteroids, and significantly better than vitamin D alone. Only 25 trials assessed clinical cutaneous dermal atrophy; few cases were detected, but trials reported insufficient information to determine whether assessment methods were robust. Clinical measurements of dermal atrophy are insensitive and detect only the most severe cases. No comparison of topical agents found a significant difference in systemic adverse effects.

Authors' conclusions

Corticosteroids perform at least as well as vitamin D analogues, and they are associated with a lower incidence of local adverse events. However, for people with chronic plaque psoriasis receiving long-term treatment with corticosteroids, there remains a lack of evidence about the risk of skin dermal atrophy. Further research is required to inform long-term maintenance treatment and provide appropriate safety data.

PLAIN LANGUAGE SUMMARY

Skin treatments for chronic plaque psoriasis

Chronic plaque psoriasis is the most common type of psoriasis. Although any part of the body may be affected, the most commonly affected sites are the elbows, knees, and scalp. 'Topical' treatments (i.e. treatments applied to the skin) are usually tried first. These include vitamin D products, topical corticosteroids, tar-based preparations, dithranol, salicylic acid, and vitamin A products. As chronic plaque psoriasis is a long-term condition, it is important to find out which treatments work best and what adverse effects they have. This review describes average benefits of different treatments, while recognising that individuals will vary in their experience of each treatment.

The evidence was based on 177 studies, which, in total, included 34,808 people. Studies were typically about 7 weeks' long, but this ranged from 1 week to 52 weeks. Vitamin D products were found to work better than placebo (the base cream or ointment). Potent topical corticosteroids (strong, e.g. betamethasone dipropionate) and very potent (very strong, e.g. clobetasol propionate) topical corticosteroids were also effective.

Some studies compared vitamin D products directly with potent or very potent corticosteroids. These products had similar effects when applied to the body, but corticosteroids worked better than vitamin D for scalp psoriasis. Treatment that combined vitamin D with a corticosteroid was more effective than vitamin D alone and more effective than the topical corticosteroid alone. Vitamin D products generally performed better than coal tar, but studies found conflicting results when comparing vitamin D with dithranol.

Whether applied to the body or to the scalp, potent corticosteroids were less likely than vitamin D to cause 'local adverse events', such as skin irritation or burning, and people were therefore more likely to stop using vitamin D products. When studies examined whether topical treatments had effects within the body ('systemic adverse events'), we found no difference between placebo and any other treatment. However, this may be because many trials did not properly assess systemic adverse events, rather than because there really was no difference.

More long-term studies would help doctors and people with psoriasis decide on the best way to treat this chronic condition.

BACKGROUND

Description of the condition

Psoriasis is a chronic inflammatory skin disease with a prevalence ranging from between 1% and 2% in the UK and northern European populations (Hellgren 1967; Krueger 1984) to 0.1% to 0.3% in the Far East (Simons 1949) and China (Yip 1984). Psoriasis comprises multiple phenotypes and may be localised (e.g. to the skin-fold areas (inverse psoriasis), the palms, or the soles) or widespread. Types of widespread psoriasis include guttate, generalised pustular, and erythrodermic (Griffiths 2007). Chronic plaque psoriasis may be localised or widespread and accounts for

90% of psoriasis cases (Griffiths 2007); it is characterised by red patches of thickened skin (plaques) covered in silver scales (Figure 1). Any area of the body may be affected, but the main areas are the knees, elbows, lower back, and scalp. There is a wide spectrum of disease severity from a single plaque to involvement of more than 90% of the skin surface. Psoriasis may be classified as 'mild', 'moderate', or 'severe', although these categories are difficult to define precisely (Krueger 2000). Psoriatic arthritis accompanies the cutaneous (skin) manifestations of psoriasis in 5% to 30% of cases (Barisic-Drusko 1994; Krueger 1984; Salvarani 1995; Zanolli 1992). Recent improvements in the classification criteria may reduce the wide variation in reported prevalence of psoriatic arthritis (Taylor 2006). Psoriasis occurs in 5% of people with Crohn's disease (Lee 1990).

Figure 1. Chronic plaque psoriasisSource: Dermis Dermatology Atlas Online (used with permission)



Causes

The way that psoriasis develops is complicated and appears to be influenced by many factors, including genetic changes, local trauma, infections, certain drugs (such as beta-blockers, lithium, chloroquine, and non-steroidal anti-inflammatory drugs (NSAIDs)), the duration of antipsoriatic treatments, endocrine factors, sunlight, alcohol, smoking, and stress (Tagami 1997). The skin lesions of psoriasis are shown in Figure 2, and they are characterised by cells multiplying too quickly (epidermal hyperproliferation), cells not maturing normally (abnormal keratinocyte differentiation), and the presence of cells that cause inflammation (a lymphocyte inflammatory infiltrate) (Barker 1991; Griffiths 2003; Stern 1997). Psoriasis is now recognised as an immune-mediated disorder, with tumour necrosis factor alpha (TNF α), dendritic cells, and T-cells all contributing to its pathogenesis (Griffiths 2007a). Several genes interact with environmental factors to induce the development of psoriasis, and different combinations of changes in several genes and environmental factors can produce the same clinical picture of psoriasis (Bhalerao 1998; Brandrup 1978; Farber 1974; Lomholt 1963; Willan 1808). A locus (plural = loci) is the specific location of a gene on a chromosome, and its

position is defined using the letters 'p' (for a chromosome's short arm) and 'q' (for a long arm). At least nine chromosomal psoriasis susceptibility loci were originally identified (Griffiths 2007a). The strongest association and linkage is to a locus within the major histocompatibility complex, the area affecting immune response (Genetic Analysis of Psoriasis Consortium 2010; Henseler 1992; Russell 1972; Svejgaard 1974; Tazi-Ahnini 1999a; Tazi-Ahnini 1999b; Trembath 1997). Other linkage studies have reported linkage to 4q and 17q (Matthews 1996; Tomfohrde 1994) and 16q and 20q (Nair 1997; Trembath 1997). Proinflammatory CD4-positive T helper cells produce interferon gamma (produced by Th1) or interleukin (IL)-17 (produced by Th17). These cells interact with dendritic cells, macrophages, mast cells, and neutrophils, causing inflammation (Ghoreschi 2007). A meta-analysis of 3 genomewide association studies (GWAS) has identified 15 new susceptibility loci (Tsoi 2012) for psoriasis. This brings the total number of loci associated with psoriasis to 36. Several of these loci are involved in the regulation of the skin's innate immune response. They provide confirmation of the role of several existing biologic therapies as well as new targets for drug development.

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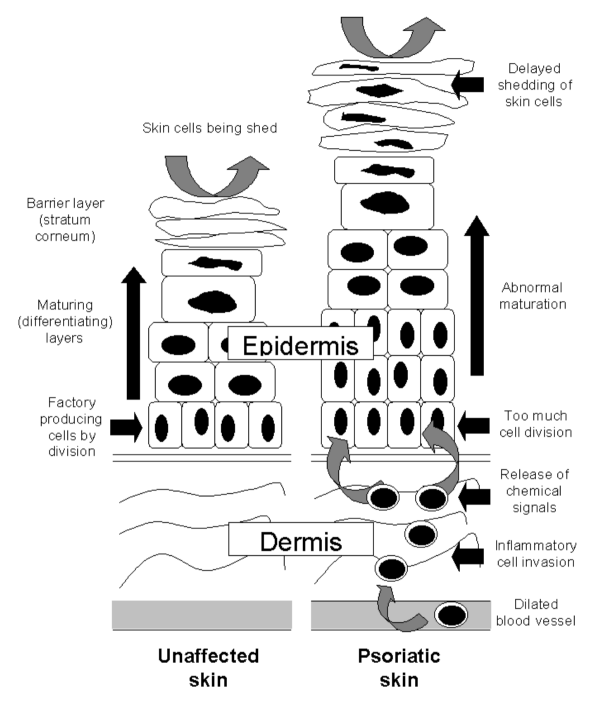


Figure 2. The epidermis in the skin of people with and without psoriasis

Impact

Until identified as a single disease by von Hebra in 1841, psoriasis was thought to be a variant of leprosy and regarded as contagious (de Jong 1997). The misconception may persist: In a survey of people with psoriasis in 1997, almost three-quarters of respondents reported that others thought their condition was contagious, and a similar proportion feared swimming and taking part in sporting activities (Watts 1998). Psoriasis can lead to social isolation (van de Kerkhof 1997a), stigmatisation (Gupta 1998; van de Kerkhof 1997a), and fear of other people's reactions, adversely affecting the quality of daily life (Finlay 1994; Finlay 1995a; Finlay 1995b; Finlay 2001; McKenna 2003; Ortonne 2000; Richards 2003; Stern 1995). Psychological distress induced by psoriasis may also impair the response to treatment (Fortune 2003).

Description of the intervention

Treatment of psoriasis should always be appropriate to its severity and importance to that individual: It should never be more unpleasant, intolerable, or dangerous than the disease itself (Camp 1992). Topical treatments include vitamin D analogues, topical corticosteroids, tar-based preparations, dithranol, salicylic acid, and topical retinoids (Baadsgaard 1995; Corbett 1976; Fredriksson 1980; Goeckerman 1931; Ingram 1953; Kragballe 1988; Kragballe 1989; Langner 1996; Staberg 1989; Unna 1916; Van de Kerkhof 1996a), but there is no evidence-based 'treatment ladder' by which to sequence treatments (Van de Kerkhof 2008). Emollients are generally used in a supportive role as an addition to topical treatments, to normalise hyperproliferation, differentiation, and to exert anti-inflammatory effects (Fluhr 2008). The two classes of topical treatment for psoriasis that are most commonly prescribed in developed countries are vitamin D analogues and topical corticosteroids, because they are considered more cosmetically acceptable than tar and dithranol preparations (Baadsgaard 1995; Kragballe 1988; Van de Kerkhof 1996a).

Topical corticosteroids (specifically glucocorticoids) are available in four potencies: mild, moderate, potent, and very potent, which are assessed using the vasoconstrictor assay (BMA 2012). The benefit of topical steroids is that in cream formulations, they are easy to apply, cosmetically acceptable, do not stain the skin, and rarely cause irritation. There are several adverse effects of corticosteroids, including cutaneous atrophy, rebound after discontinuation of treatment, and decreasing response to the drug (tachyphylaxis) (du Vivier 1975; Lee 1998; Kao 2003). Glucocorticoids (GC) exert their effects either via interaction with cell membranes (non-genomic effects) or downstream with the genome and via interaction with intracellular fluid in GC receptors and downstream with the genome (genomic effects). The genomic effects are of two types: "transrepression (inhibition of synthesis of regulatory proteins) and transactivation (induction of the synthesis of regulatory proteins)" (Bos 2008). Transactivation appears to mediate certain adverse reactions, such as cutaneous atrophy. Immunomodulation seems to be the result of GC-mediated transrepression, that is, silencing of proinflammatory genes, such as TNF α . Non-steroidal GC receptor ligands (selective GC receptor agonists) have recently been identified and may reduce the side-effects of GC without loss of immunosuppressive effects (Bos 2008).

The naturally occurring active metabolite of vitamin D, calcitriol (1a,25-dihydroxyvitamin D3) (Langner 1996), and two synthetic vitamin D analogues, calcipotriol (Kragballe 1988; Kragballe 1989; Staberg 1989) and tacalcitol (1a,24-dihydroxyvitamin D3) (Baadsgaard 1995; Van de Kerkhof 1996a), are effective when applied topically in psoriasis (Mason 2002a). These agents bind to vitamin D receptors (VDR), which in turn bind to vitamin Dresponsive elements (VDRE) in multiple genes. 'Switching on' (transactivation of) these genes inhibits the multiplication of cells and stimulates their differentiation (Figure 2). VDRs also suppress the inflammatory component of psoriasis by inhibiting the production of proinflammatory cytokines (small proteins that affect cell-cell interaction), such as interleukin-1 (IL-1). Vitamin D analogues all have the potential to induce abnormally high levels of calcium in the blood serum (hypercalcaemia) and urine (hypercalciuria). Although calcipotriol ointment causes no elevation of total serum calcium when used at the recommended dose of 100 g per week (Mortensen 1993), there are significant elevations in both serum and urinary calcium when the dose is increased to 300 g per week (Bourke 1993a; Bourke 1994). Topical vitamin D analogues are cosmetically acceptable; they are not known to cause skin atrophy; and they are not usually associated with rebound when therapy is discontinued. However, at least 25% of people are reported to have little or no response to topical vitamin D analogues (Holick 1996; Mee 1998).

Urea or salicylic acid may be used to reduce thickness and scaling of the skin; combination with other products can improve their absorption. However, these can also irritate the skin. Topical immunosuppressants, such as methotrexate, and topical macrolactams, such as tacrolimus, are relatively new treatments, and their effectiveness, tolerability, and longer-term effects are less clear than with the more established products. This review also considers combination products involving any of the above treatments.

The Cochrane Library has three published Cochrane reviews of interventions for psoriasis. Owen 2000 assessed the impact of antistreptococcal interventions for guttate and chronic plaque psoriasis. The review found that "although both antibiotics and tonsillectomy have frequently been advocated for patients with recurrent guttate psoriasis or chronic plaque psoriasis, there is to date no good evidence that either intervention is beneficial." Chalmers 2000 reviewed all treatments, excluding antistreptococcal interventions, for guttate psoriasis. The review identified only one relevant trial and no evidence of the effectiveness of any topical interventions. Chalmers 2006 assessed interventions, including topical treatments, for chronic palmoplantar pustulosis (a disease that is closely related to psoriasis and used to be considered a variant of psoriasis). Chalmers 2006 found that topical steroids under hydrocolloid occlusion were effective in inducing remission. In addition, *The Cochrane Library* has four published Cochrane review protocols, which cover interventions for nail psoriasis (Velema 2009), interventions for scalp psoriasis (Jales 2012), phototherapy (Chen 2011), and the biological agent ustekinumab (Roberts 2011).

Why it is important to do this review

Chronic plaque psoriasis is a condition for which there is no known cure, and currently, available treatments may only temporarily clear the skin (Bonifati 1998; Griffiths 2004). Clinical practice varies between and within different countries. By focusing on topical treatments for psoriasis, either as monotherapy or in combination, this review assesses the relative effectiveness, tolerability, and safety of these treatments and so helps to determine how best to induce remission and delay recurrence in people receiving topical treatment. Table 1 provides a list of acronyms used in the review.

Structure of the Review

The structure of the review is provided to facilitate navigation:

- Objectives
- Methods
- Results
- • Description of the studies
 - o Risk of bias in the included studies
 - Effects of the interventions
 - \diamond (1) Primary outcome measures

(a) Investigator's Assessment of Overall Global Improvement (IAGI)/Investigator's Global Assessment of Disease Severity (IGA)

- (b) Total Severity Scores (TSS)
- (c) Psoriasis Area and Severity Index (PASI)

(d) Patient Assessment of overall Global Improvement (PAGI)/ Patient Global Assessment of Disease Severity (PGA)

(e) Combined end point (IAGI/TSS/PASI/PAGI)

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♦ (2) Secondary outcome measures
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(a) Withdrawal rates (total rate; withdrawal because of adverse events; withdrawal because of treatment failure)

(b) Adverse events (local and systemic)

(i) Findings from the main review

(ii) Findings from the separate search for additional studies of adverse events

- (c) Quality of life measures
- (d) Economic outcomes (not updated in 2011)

(e) Concordance or adherence with treatment (not updated in 2011)

- Discussion
- Authors' conclusions

Under 'Primary outcome measures', we report findings for each of the 19 analyses (including sensitivity analyses). We also do this under 'Secondary outcome measures' for subsections (a) and (b). We did not update the sections on Economic outcomes (2d) and Concordance (2e) in 2011 because of resource constraints.

OBJECTIVES

To compare the effectiveness, tolerability, and safety of topical treatments for chronic plaque psoriasis, relative to placebo, and to similarly compare vitamin D analogues (used alone or in combination) with other topical treatments.

METHODS

Criteria for considering studies for this review

Types of studies

We included randomised controlled trials in the review. Trials could be either placebo-controlled or head-to-head with a vitamin D preparation (head-to-head trials compare two active treatments with each other). The types of study design eligible for inclusion were as follows: parallel-group (between-patient), cross-over, and within-patient designs. For within-patient studies, where study participants serve as their own control, we included only those studies that clearly adopted a left-right design, and we excluded studies where multiple plaques were treated with more than two products. If no useful effectiveness, withdrawal, or adverse events data were available, either from the published paper or from sponsors or trialists, we excluded the study.

In addition to findings on adverse events from the main review, we undertook separate searches for additional safety and tolerability studies. The searches for longer-term adverse events included studies of any design that included humans (i.e. not only animals; either humans only or humans and animals). However, studies with fewer than 10 participants (including case reports) were not eligible for inclusion. We did not restrict the search for concordance/ adherence studies by study design (i.e. non-randomised studies were eligible for inclusion).

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Types of participants

People of any age with chronic plaque psoriasis affecting the body, limbs, scalp, or a combination of the aforementioned. We did not limit participant type by area of involvement, disease severity, or skin area treated.

Types of interventions

Topical treatments, including the following:

- vitamin D preparations, e.g. calcipotriol;
- corticosteroids, e.g. betamethasone valerate;
- coal tar;
- dithranol, also known as anthralin;
- salicylic acid;
- urea;
- topical retinoids;
- topical immunosuppressants, e.g. methotrexate;
- topical macrolactams, e.g. ascomycin derivatives, such as tacrolimus; and

• combination products, e.g. corticosteroids with coal tar or corticosteroids with vitamin D.

We compared topical treatments with vehicle (placebo). We also compared vitamin D analogues with other topical treatments. We selected vitamin D analogues for this comparison because they are first-line treatments in many developed countries (van de Kerkhof 1998). We based the potency of topical corticosteroids on classifications from a previous review (Mason 2002b).

The review included any topical treatment for psoriasis, except for products for which (a) no licence was obtained and (b) research into the product was discontinued. The reason for this exclusion criterion is that these products are unlikely to be of interest to people making decisions about health care, such as policy-makers, people with psoriasis, or clinicians. Although they may be of interest to researchers, lessons from the research into 'failed' molecules are likely to have been reflected in the development of subsequent products.

Trials of systemic or ultraviolet (UV) (phototherapy) treatments with adjunctive topical treatment were not eligible for inclusion in the review.

Types of outcome measures

Table 2 provides an overview of the effectiveness outcome measures included in the review. We provide details of how we used the primary outcomes to derive a 'combined end point' in the section 'Measures of treatment effect'.

Primary outcomes

1. Investigator's Assessment of Overall Global Improvement (IAGI)/Investigator's Global Assessment of Disease Severity (IGA).

- 2. Total Severity Scores (TSS).
- 3. Psoriasis Area and Severity Index (PASI).

4. Patient Assessment of overall Global Improvement (PAGI)/

Patient Global Assessment of Disease Severity (PGA).

Secondary outcomes

1. Withdrawal rates (total rate; withdrawal due to adverse events; withdrawal due to treatment failure).

- 2. Adverse events (local and systemic).
- 3. Quality of life measures.
- 4. Economic outcomes.
- 5. Concordance or adherence with treatment.

Search methods for identification of studies

We aimed to identify all relevant randomised controlled trials (RCTs) regardless of language or publication status (published, unpublished, in press, or in progress).

Electronic searches

Search strategies used for the previous version of the review (Mason 2009; see also Acknowledgements) were revised where appropriate and rerun. We did not restrict the searches by body area affected. The information specialists updated the search strategies to reflect changes in the interfaces and MeSH (Medical Subject) headings, as well as to incorporate terms for newly licensed products. In February 2011, the following databases were searched for effectiveness RCTs of psoriasis treatments:

• the Cochrane Skin Group Specialised Skin Register (searched 8 February 2011) using the search strategy in Appendix 1;

• the Cochrane Central Register of Controlled Trials (CENTRAL) in *The Cochrane Library* (2011, Issue 2) using the search strategy in Appendix 2;

• MEDLINE via OVID (from 1948) using the strategy in Appendix 3;

• EMBASE via OVID (from 1980) using the strategy in Appendix 4;

• Science Citation Index (SCI) via the Institute for Scientific Information (ISI) Web of Knowledge interface (now known as Thomson Reuters) (from 2008) using the strategy in Appendix 5;

• Conference Proceedings Citation Index - Science (CPCI-S) via the ISI web of Knowledge interface (from 2008) using the strategy in Appendix 5;

• BIOSIS via the DialogClassic interface (from 1993) using the strategy in Appendix 6;

• Dissertation Abstracts via DialogClassic interface (from inception) using the strategy in Appendix 7;

• Inside Conferences via DialogClassic interface (from inception) using the strategy in Appendix 7;

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• System for Information on Grey Literature in Europe (SIGLE) via WebSPIRS interface (search not updated) using the strategy in Appendix 8;

• National Research Register (NRR) (CD-ROM interface, issue 2004/4) using the strategy in Appendix 2; and

• the UK Clinical Research Network Study Portfolio (http://public.ukcrn.org.uk/search/) using the strategy in Appendix 2.

To comply with Cochrane policy (stipulating that reviews must be published within 12 months of the electronic searches being run), further searches for this update were run on 23, 24, and 29 August 2012. Although it was not possible to incorporate RCTs identified through this search within this review, we listed relevant references in the 'Characteristics of studies awaiting classification' tables. They will be incorporated into the next update of the review.

Searching other resources

References from published studies and reviews

We checked these for further references to relevant trials.

Unpublished literature

We routinely contacted trialists and companies for newly published studies and missing data.

The *meta*Register of Controlled Trials (http://www.controlledtrials.com/mrct/) was searched in August 2012 for ongoing and unpublished trials.

Adverse effects

On 2 February 2011, the following databases were searched for studies of adverse events of specific psoriasis treatments:

- MEDLINE via OVID (see Appendix 9); and
- EMBASE via OVID (see Appendix 10).

We limited searches to English-language papers published in the years between 2005 to 2011. In MEDLINE, the search was designed to omit records with the following publication types: 'note', 'comment', and 'editorial'.

We also considered relevant adverse effects studies identified during the screening for effectiveness trials.

These searches were updated in August 2012, identifying 537 new references. We will incorporate these studies into the next update of this review.

Concordance/adherence

We did not undertake searches for concordance/adherence in the 2011 review update because of resource constraints.

Language restrictions

There were no language restrictions when searching for effectiveness RCTs or concordance/adherence studies. We restricted searches for studies of adverse events to those published in English.

Data collection and analysis

Selection of studies

Two authors (AM and JM) screened titles and (where available) abstracts identified from the searches, and another author (MC) acted as an arbiter when necessary. In our protocol, we stated our intention that we would exclude studies meeting only some of the inclusion criteria stated above. However, this was infeasible, because we would have needed to cite large numbers of studies (over 1000). Therefore, we listed as excluded only those studies that we deemed potentially eligible for inclusion *and* for which we retrieved full papers, but which subsequently failed to meet the inclusion criteria.

For the separate search for studies exploring adverse events, we deemed studies as eligible if they addressed safety or tolerability issues, focused on drugs included in the main review, and were longer-term in follow-up (> 12 weeks). Short-term studies (with follow-up < 12 weeks) were eligible for inclusion only if they were designed specifically to consider adverse effects, tolerability, or safety. Studies that included fewer than 10 participants (including case reports) were not eligible for inclusion.

For the separate search for studies of concordance/adherence with treatment, studies were eligible if they addressed adherence with topical treatment in people with any type of psoriasis. This section was not updated because of resource constraints.

Data extraction and management

Applying methods from our original review (Mason 2002a), we summarised the major attributes of trials, including treatment forms, doses and duration, inclusion and exclusion criteria, level of blinding, within-patient or between-patient (parallel-group) design, method of generation of the randomisation sequence, concealment of allocation, numbers of participants randomised, baseline comparability, loss to follow up, primary and secondary outcomes, withdrawals, and adverse events. One reviewer (AM) extracted the data, and another reviewer (HH) checked these data. We extracted data from trials on four primary outcomes:

1. IAGI (Investigator's Assessment of Global Improvement) or the IGA (Investigator's Global Assessment of Disease Severity).

- 2. TSS (Total Severity Score).
- 3. PASI (Psoriasis Area and Severity Index).
- 4. PAGI (Patient Assessment of Global Improvement) or the PGA (Patient Global Assessment of Disease Severity).

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Where available, we also extracted data on quality of life, economic outcomes, and concordance/adherence.

In addition, we extracted data on withdrawal due to any reason, such as adverse events or treatment failure, as well as adverse events due to local and systemic effects.

For each outcome measure under a comparison, we included the same treatment options regardless of data availability (see Data and analyses). We did this for three reasons. Firstly, an inclusive approach makes clear that there is an absence of data, not that data have been omitted. Secondly, if data subsequently become available when the review is updated in future, the correct structure is in place for data entry. Thirdly, this approach ensures treatments are always ordered identically regardless of outcome.

Assessment of risk of bias in included studies

Assessment of methodological quality

The quality assessment included an evaluation of each included study, based on the following components, which are considered to be associated with biased estimates of treatment effect (Juni 2001):

(a) the method of generation of the randomisation sequence;

(b) the method of allocation concealment - we considered this 'adequate' if the assignment could not be foreseen;

(c) who was blinded/not blinded (participants, clinicians, outcome assessors); and

(d) how many participants were lost to follow up.

In addition, the quality assessment included the following:

(e) baseline assessment of the participants for age, sex, duration, and severity of psoriasis; and

(f) baseline comparability of intervention and control groups.

We recorded the information in the 'Characteristics of included studies' section.

Measures of treatment effect

Summarising primary outcomes with standardised mean differences

We extracted data on four primary outcome measures:

• IAGI (Investigator's Assessment of Global Improvement) or the IGA (Investigator's Global Assessment of Disease Severity)

- TSS (Total Severity Score)
- PASI (Psoriasis Area and Severity Index)

• PAGI (Patient Assessment of Global Improvement) or the PGA (Patient Global Assessment of Disease Severity)

Trials often reported more than one measure, but none of the trials reported all measures. We therefore devised a 'combined

end point', which allowed more data to contribute to an overall analysis and facilitated treatment comparisons. We labelled this 'super' outcome as outcome (e) throughout the review.

We constructed the combined end point by taking IAGI (or IGA) data when available, and failing this, TSS, PASI, or PAGI (PGA) data in that order of availability. For PASI and TSS, some included trials reported change scores and others reported end point scores. In view of the mix of end point/change scores and of the variation in scale, we analysed findings using a standardised mean difference statistic (SMD) in a random-effects model. Table 2 summarises the characteristics of the outcome measures.

We also expressed SMDs in physical units adjusting by the appropriate pooled standard deviation estimate (Table 3).

Secondary outcomes

We summarised data on adverse events, quality of life measures, economic outcomes, and concordance as narratives. We summarised withdrawal data using the risk difference (RD) metric and pooled using a random-effects model. We felt this was more appropriate than a fixed-effect model since definitions of withdrawal and adverse events vary between trials.

Unit of analysis issues

Within-patient studies are statistically analogous to cross-over studies, and results should be adjusted by the correlation coefficient (Section 16.4.6, *Cochrane Handbook for Systematic Reviews of Interventions*; Higgins 2011). No study included in the review reported this statistic, and we did not have access to patient-level data, so could not estimate it directly.

On the subject of cross-over studies, the Cochrane Handbook for Systematic Reviews of Interventions states (Section 16.4.5; Higgins 2011) the following: "A common situation is that means and standard deviations (or standard errors) are available only for measurements on E [experimental group] and C [control group] separately. A simple approach to incorporating cross-over trials in a meta-analysis is thus to take all measurements from intervention E periods and all measurements from intervention C periods and analyse these as if the trial were a parallel-group trial of E versus C. This approach gives rise to a unit-of-analysis error (see Chapter 9, Section 9.3) and should be avoided unless it can be demonstrated that the results approximate those from a paired analysis, as described in Section 16.4.4. The reason for this is that confidence intervals are likely to be too wide, and the trial will receive too little weight, with the possible consequence of disguising clinically important heterogeneity. Nevertheless, this incorrect analysis is conservative, in that studies are under-weighted rather than overweighted. While some argue against the inclusion of cross-over trials in this way, the unit-of-analysis error might be regarded as less serious than some other types of unit-of-analysis error."

Consequently, we included within-patient studies as though they were parallel-group studies, accepting that they are under-

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weighted. To explore whether it was appropriate to combine these trials, we undertook two sensitivity analyses. First, we considered how effect size varied for within- and between-patient studies. If the magnitude of effect varied consistently between the two study designs, this strongly suggested a non-zero correlation coefficient and an appropriateness to separate the trials. Second, we used sensitivity analysis to explore the impact on pooled findings of varying the correlation coefficient (rho) for within-patient studies. This analysis used the generic inverse variance measure, with SMDs and their standard errors estimated from the formulae in the *Cochrane Handbook for Systematic Reviews of Interventions* (Section 16.4.6.4) (Higgins 2011). These estimated SMDs differ slightly from those that RevMan estimates for continuous outcomes, even when the correlation coefficient is zero (which is the assumption implicit in the latter model).

The analyses found no evidence that the magnitude of effect varied consistently. Within-patient trials did not consistently demonstrate smaller or larger effects than between-patient trials. Varying the value of rho had no significant effect on the findings: As rho increased, the effect size increased and the confidence intervals (usually) widened, but the magnitudes of changes were small and non-significant at the 5% level.

In the interests of statistical purity, these trials could (a) be reported separately or (b) be removed altogether. The drawback of option (a) is that it makes an already complex review even more complex and less accessible; the disadvantage of option (b) is that it removes data that might be of interest to clinicians and people with psoriasis. On balance, we preferred to report relevant randomised data wherever possible to help inform pragmatic decision-making.

Dealing with missing data

We routinely contacted trialists and companies for missing data. Where studies did not report estimates of variance, we derived them from confidence intervals (CIs) or from P values where possible. Where we could not obtain estimates of variance, we imputed them deterministically by pooling the standard deviations of treatment cohorts fully reported in trials and adjusted for scale size.

We made separate imputations for each outcome measure (see Table 3):

- for within-patient studies;
- for between-patient (parallel-group) studies;
- for end point scores;
- for change scores; and
- for scalp trials.

Within-patient designs are statistically analogous to cross-over studies, and the precision of their findings within a meta-analysis needs adjustment for within-patient correlation. We attempted to explore this by sensitivity analysis.

Data synthesis

We analysed findings using a standardised mean difference statistic (SMD) in a random-effects model. However, this model cannot perfectly address all the sources of design complexity that arise when summarising findings across studies. Three of the main sources of complexity are listed below. Other sources of complexity include variation in trial duration, disease severity, participant demographics, treatment application method, dosing frequency, drug potency and vehicle.

Study design (within- versus between-patient)

Trials were either between-patient or within-patient designs. The former randomise participants into separate (parallel) groups; the latter randomise treatments to the left or right side of the same participant. Within- and between-patient trials have different variance structures. Moreover, the two responses (left and right) of within-patient studies may be correlated (See Unit of analysis issues for further details).

Absence of a simple one-to-one correspondence between papers, trials, and comparisons

There were instances of single papers reporting either multiple trials or multiple analyses within a single trial. Therefore, simple counts of numbers of participants and numbers of studies contributing data to the analysis were misleading, and we made adjustments accordingly (Table 4). Thus, these numbers may not match the numbers estimated in RevMan, which does not account for these factors.

Body area targeted for treatment

Whereas the majority of trials investigated chronic plaque psoriasis on the body, some trials focused on scalp psoriasis; some reported findings for both scalp and body psoriasis; and some were of inverse (flexural) or facial psoriasis. One trial of body and scalp psoriasis reported overall outcomes (IAGI/PAGI), a scalponly outcome (TSS), and a body-only outcome (modified PASI) (Van de Kerkhof 2002a). Ortonne 2010 reported findings separately for treatment of the body and treatment of the face. We previously used sensitivity analysis to investigate the scalp psoriasis trials (Mason 2009), but in this update we analysed trials of inverse psoriasis (comparisons 16 and 17) and scalp trials (comparisons 18 and 19) separately from the trials of body psoriasis (see Sensitivity analysis).

Subgroup analysis and investigation of heterogeneity

We examined findings by agent class (as our primary analysis) and individual topical agent (within-class analysis).

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When comparing trials both within and across therapeutic classes, the summary estimates may demonstrate substantial heterogeneity. Ideally, we would seek to identify the reasons for individual differences, but publications rarely report sufficient detail to make a robust investigation feasible. Reasons might include differences in trial design, length of follow-up, disease severity, participant selection, adherence, adequacy of concealment of allocation, adequacy of blinding, and source of funding (Mason 2002a).

The Cochrane Handbook for Systematic Reviews of Interventions explicitly endorses the combination of 'apples and oranges' "if they are used to contribute to a wider question about fruit" (Section 9.5.1; Higgins 2011). Our purpose was to identify whether classes of topical treatments work and are safe. To this end, there is a fundamental difference between heterogeneity that makes it uncertain whether individual people with psoriasis will derive any benefit from a treatment and heterogeneity that makes the size of a positive benefit imprecise. Clinicians and those with psoriasis will still value information about a treatment that is beneficial even though its magnitude is poorly understood. However, we clearly stated the presence of heterogeneity where it occurred and used the Cochrane Handbook for Systematic Reviews of Interventions as a guide to interpretation (Section 8.5.2; Higgins 2011).

Sensitivity analysis

We used a meta-analysis with a random-effects estimation both for measures of effect and for pooling of risk differences for adverse events. We quantified heterogeneity using the I² statistic. If we identified outliers, we undertook a sensitivity analysis to investigate the implications of their exclusion on the pooled summary statistics. In addition, we undertook sensitivity analyses to investigate the impact of within-patient versus between-patient trials, and to explore the impact on pooled findings of varying the correlation coefficient (see Unit of analysis issues). In some comparisons, there were no, or relatively few, studies that included both within-patient and between-patient designs, few participants contributing data, or both. We used the following criteria to help decide whether we should have included an analysis in the sensitivity analysis:

• frequently-used products in clinical practice; and

• for within-/between-patient sensitivity analysis: whether it included both within-patient and between-patient designs

Where at least two within-patient trials were included in a pooled comparison, we explored the potential influence of the correlation coefficient.

Based on these criteria, we selected six comparisons (analyses 1, 2, 3, 4, 7, and 18) for sensitivity analysis. To ensure sufficient data were available, we analysed the combined end points. These analyses cover vitamin D analogues, dithranol, and corticosteroids, which are amongst the most frequently used products in clinical practice.

Other

We involved a consumer throughout the review process to help ensure the readability of the final review.

RESULTS

Description of studies

Results of the search

For this update, the RCT searches identified 3749 records:

- MEDLINE: 1312
- EMBASE: 2008
- SCI: 253
- BIOSIS: 44
- Dissertation Abstracts: 1
- Inside Conferences: 0
- CENTRAL: 70
- UK Clinical Research Network: 20
- Skin Group Specialised Register: 41

The total number of new records assessed after deduplication against each other and previously identified records was 2637. We added records from the searches in February 2011 to those identified from searches run in 2008 (see Mason 2009). The total number of records screened for this review over time is now 5414. From the 2011 searches, we retrieved 148 papers and screened these for eligibility. (Some papers were multiple reports of the same trial).

In 40 trials (some of which were consequently excluded), some or all outcome data were missing. We contacted trialists or sponsors to request missing data, receiving data for 25 of these trials. We excluded trials that reported no useable outcome data. We did not contact trialists or sponsors for missing adverse events or withdrawal data, although some sponsors provided this spontaneously. We included 48 new randomised controlled trials in the updated review. Compared with the previous version of this review (Mason 2009), studies were larger (mean number of participants: 284 versus 164), had a longer treatment duration (10 weeks versus 6 weeks) and follow up (11 weeks versus 8 weeks), and were more likely to be parallel-group in design (88% versus 63%). The new studies were also more likely to include an active control group (60% versus 43%) and patient-reported outcomes (44% versus 24%), and there were relatively more scalp trials (21% versus 12%). The 48 trials provided evidence on 7 new active treatments.

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Included studies

The updated review included 177 studies, with 34,808 participants.

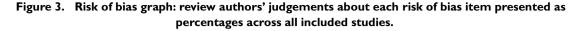
The number of included studies counted by RevMan is 190, because we entered each study reporting a placebo and an active comparison into the 'Characteristics of included studies' table as two studies.

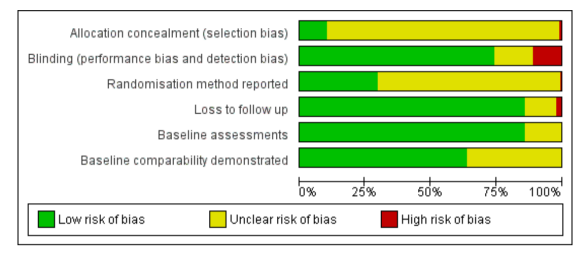
Of the included studies, 106 of these were placebo-controlled; 84 compared treatments head-to-head, with 15 trials reporting both placebo-controlled and head-to-head comparisons. The 15 trials reporting both head-to-head and placebo comparisons contributed only once to the analysis of study characteristics, unless the trial involved entirely distinct participants in its placebo-controlled and active-controlled analyses. For example, the trial by Guenther 2002 compared treatments against each other (Guenther 2002 (H)) and against placebo (Guenther 2002 (P)). This study contributed only once to the analysis of study characteristics (number of participants, proportion of males, etc). However, two trials reported placebo and head-to-head analyses involving entirely separate participants (Barker 1999 (H) and Barker 1999 (P); Grattan 1997 (H) and Grattan 1997 (P)). Therefore, the total number of studies contributing data to the analysis of study characteristics and quality assessment was 177 (106 placebo + 84 head-to-head -15 double-counted trials (with placebo and active comparators) and 2 trials that each report 2 separate studies (Barker 1999 (H) and Barker 1999 (P); Grattan 1997 (H) and Grattan 1997 (P)). There were 26 trials of scalp psoriasis (Barrett 2005; Buckley 2008; Cook-Bolden 2010; Duweb 2000; Elie 1983; Ellis 1988; Franz 1999; Franz 2000; Green 1994; Jarratt 2004; Jemec 2008 (H) and Jemec 2008 (P); Kiss 1996; Klaber 1994; Klaber 2000b; Köse 1997; Kragballe 2009; Lepaw 1978; Luger 2008; Olsen 1991; Pauporte 2004; Poulin 2010; Reygagne 2005; Shuttleworth 1998; Tyring 2010; Van de Kerkhof 2002a; Van de Kerkhof 2009). Six trials investigated inverse psoriasis, facial psoriasis, or both (Gribetz 2004; Kreuter 2006 (H) and Kreuter 2006 (P); Lebwohl 2004; Liao 2007; Ortonne 2003; Ortonne 2010). One trial evaluated psoriasis in children (Oranje 1997). Most trials were conducted in ambulatory care settings, but four trials were of hospitalised participants (Grattan 1997 (H) and Grattan 1997 (P); Kragballe 1991a; Monastirli 2000; Van der Vleuten 1995).

One hundred and twenty-three trials adopted a between-patient (parallel-group) design; 53 were within-patient studies; and one trial used both designs (Henneicke-v. Z. 1993). The trial by Levine (Levine 2010 (H) and Levine 2010 (P)) was a within-patient trial that randomised participants to two of seven treatment options. Therefore, the pair-wise comparisons we analysed (e.g. calcipotriol versus placebo) included a mixture of within- and between-patient designs: Some participants received calcipotriol on one side and placebo on the other; other participants received calcipotriol on one side and another active treatment on the other side.

The 177 studies included 34,808 participants. Of these studies, 133 provided data on the age of participants. The mean age of all participants for which studies provided data was 47.2 years (range = 2 to 97 years) (N = 28,921). Data on the gender of participants (N = 28,941) were available from 140 studies. Overall, participants were more likely to be male; the mean proportion of males was 56.7% (range = 30% to 100%).

Almost half the studies (77/177 = 44%) did not clearly report the overall baseline severity of study participants (e.g. participants with mild to moderate disease) (Figure 3). One hundred studies explicitly reported baseline severity or reported sufficient information on global severity scores, such as the mean and variation in baseline PASI or the percentage of body surface area (BSA) affected, to allow us to infer global severity using guidance on the interpretation of severity scores (Finlay 2005; Krueger 2000). In the 100 trials where severity was assessable, we classified participant severity as mild (5 studies), mild to moderate (36 studies), mild to severe (6) or very severe (2 studies), moderate (12 studies), moderate to severe (27 studies), moderately severe (8 studies), moderately severe to very severe (2 studies), and severe (8 studies).





Seventy-seven studies provided insufficient information to allow an assessment of clinical severity to be made; we could not make assessments of the clinical characteristics of participants in studies reporting only the mean PASI (with no information about variation) or reporting only localised (e.g. TSS) scores. One example of a study that included participants with a wide range of severity scores is the trial by Cunliffe 1992, where the mean baseline PASI was 9.0 (suggesting moderately severe disease, according to Finlay 2005), but where individual participant scores ranged from 0.6 to 41.2. Another example is the study by Olsen 1996 (1), where participant BSA involvement averaged 12%, but ranged from 1% to 80%. It is unclear how participant severity was distributed within these ranges (i.e. whether these extremes were 'outliers' or whether a sizeable proportion of participants were clustered at the extreme ends of the distribution).

Even where trialists classified participant severity, it was not always clear that this was consistent with published guidance, which itself does not always provide consistent messages. For example, Finlay 2005 states that a PASI score > 10, a BSA involvement > 10%, or Dermatology Life Quality Index (DLQI) score > 10 constitutes severe disease. However, Krueger 2000 argues that BSA is unreliable as an indicator of severity, which is better proxied by quality of life assessments. However, the included studies rarely assessed quality of life. Given this lack of clarity and the absence of adequate severity data in around half (44%) of the included studies, we could not use sensitivity analysis to investigate the impact of baseline participant severity, nor could we reliably use severity to investigate inter-study heterogeneity.

All 177 studies provided data on treatment duration (mean: 7 weeks; range = 1 to 52) and follow-up duration (mean: 9 weeks; range = 2 to 52), where 'follow-up duration' was defined as including the treatment period. Commonly used outcomes assessed

by the studies included the following:

• individual signs (erythema, scaling, induration) (105/177 studies = 59%);

• Total Severity Score, Total Sign Score, or equivalent (83 studies = 47%);

- PASI (65 studies = 37%);
- IAGI/IGA (113 studies = 64%); and
- PAGI/PGA (52 studies = 29%).

Outcome measures employed by small numbers (< 5) of trials included the following;

• Local Psoriasis Severity Index (scale not reported);

• Jacoby assessment score (0 to 7 score transformed to % clinical improvement); and

• investigator assessment of skin staining.

Trials seldom assessed quality of life (9 trials = 5%). Participant-reported outcomes included the following:

- overall participant assessment (relative efficacy, speed of response, irritation, staining, ease of application);
- participant global assessment of acceptability of treatment, participant assessment of likely adherence; and
 - participant assessment of cosmetic acceptability.

We grouped placebo-controlled trials by type of treatment (e.g. vitamin D products) and grouped head-to-head trials in a similar way (e.g. vitamin D versus potent corticosteroid). We included 19 comparisons in the review. Since many trials did not specify participants' disease severity, it was not possible to use severity to inform pooling decisions. The primary analysis explored the results of pooling within these 19 comparison groups using a random-effects model. In addition, we undertook sensitivity analyses for five

comparisons using the 'combined end point'. These analyses used pooled data to explore within- and between-patient trial variation. In 90% (159/177) of the studies included in the review, participants applied their own treatments. Nurses applied treatments in 1 trial (Geilen 2000); participants' parents delivered some care in a trial of childhood psoriasis (Oranje 1997); and the delivery method was unclear in 16 studies.

Excluded studies

We excluded 43 studies, of which we had newly added 16 studies in this update of the review (see 'Characteristics of excluded studies' tables). The most common reasons for exclusion from the update were that the study did not report adequate data and requests for missing data from trialists or sponsors were unsuccessful (N = 5), or that the study did not provide a comparison of interest (N = 6). Two studies were not randomised (Kaur 2004; Vena 2005); one study assessed multiple plaques (Buder 2010); and two evaluated unlicensed products that were not subsequently marketed (Agrawal 2010; Rhemus 2006). We also excluded trials of nail psoriasis that we had previously included (Mason 2009), as the topic is now covered by a separate Cochrane review (de Vries 2013).

Studies awaiting classification

Update searches were run on 23, 24, and 29 August 2012. For each database searched, we have shown below the numbers of records identified. The total number of new records assessed (after deduplication against each other and previously identified records) was 1865. Relevant studies from these searches (10 references) are listed in the Studies awaiting classification section, but we did not include them in the main review.

- MEDLINE: 1203
- EMBASE: 1140
- SCI: 129
- BIOSIS: 37
- Dissertation Abstracts: 1
- Inside Conferences: 0
- CENTRAL: 26
- UK Clinical Research Network: 0
- Skin Group Specialised Register: 67

Ongoing studies

The *meta*Register of Controlled Trials was searched for ongoing and unpublished trials during the final searches for this review in August 2012 http://www.controlled-trials.com/mrct/ using the following phrases:

• "psor* AND topical NOT completed", which retrieved 127 hits;

• "psor* AND calcipot% NOT completed", which retrieved 7 hits;

• "psor* AND vitamin D NOT completed", which retrieved 21 hits;

- "(psor* AND topical AND corticost%) NOT completed", which retrieved 63 hits; and
 - "psor* AND tar NOT completed", which retrieved 8 hits.

In total, we identified 10 potentially relevant trials. We provide details in the 'Characteristics of ongoing studies' tables.

Risk of bias in included studies

We extracted and tabulated data on six quality indicators. Summary findings are presented narratively, with characteristics for all studies presented in the 'Characteristics of included studies' tables. Figure 3 is a graphical representation of the overview of the risk of bias. All included trials were randomised, but only 47/177 (27%) clearly reported the method used to randomise participants. Concealment of treatment allocation was explicitly adequate in 15 trials, but most trials (151/177 = 85%) blinded participants to treatment allocation. Most (164/77 = 93%) trials reported loss to follow up data, and 142 trials (80%) demonstrated that groups were comparable at baseline.

Allocation

Of the 177 studies assessed for quality, 15 (8.5%) explicitly achieved adequate concealment of treatment allocation (low risk of bias). Concealment was unclear in the majority of studies (160 studies = 90.4%), so the risk of bias was also unclear. Concealment was inadequate in 2 studies (1.1%) (high risk of bias) (Figure 3).

Blinding

Most (131/177 = 74%) studies were double-blind, with 20 studies adopting a single-blind (investigator-only) approach. Eighteen studies were 'open' (no blinding), and in the remaining 8 studies, the blinding approach adopted was unclear (Figure 3). Twenty trials explicitly stated that the outcome assessor was blinded to treatment allocation. However, the outcome assessor will also have been blinded in double-blind trials where the investigator also assessed outcomes.

Incomplete outcome data

We defined 'loss to follow up' as the number of enrolled participants who failed to contribute data for the analysis.

Of the 177 studies assessed for quality, 13 (7%) provided no data on loss to follow up. For the remaining 164 studies, the mean percentage loss to follow up was 6.1% (range = 0% to 31.5%). Fifty-one studies reported that there was no loss to follow up. Four studies lost more than 25% of their participants to follow-up, and we classified them as having high risk of bias for this dimension (Henneicke-v. Z. 1993; Lin 2007; Maier 2004; Weinstein 2003) (see Figure 3).

Topical treatments for chronic plaque psoriasis (Review)

Where studies did not report estimates of variance, we derived them from confidence intervals (CIs) or from P values where possible. Where we could not obtain estimates of variance, we imputed them (see Table 3). In total, we imputed estimates of variance for at least 1 outcome measure in 45 studies (7 of which were scalp trials); details are in the notes section of the 'Characteristics of included studies' tables.

Other potential sources of bias

Method of generation of the randomisation sequence

Only randomised controlled trials were eligible for inclusion in the review. However, 130 studies (73%) did not clearly report the randomisation method used. Fourteen studies reported a block randomisation design, and 27 studies used computerised methods (5 studies used both). Four reported that sequential allocation had been used (3 of these studies were published in the 1970s); 1 study used the toss of a coin; and another study used a sealed envelope method. It could be argued that we should have excluded trials with sequential allocation from the review, but this might discriminate against studies with better reporting methods in favour of those not stating the randomisation method.

Baseline assessment of the participants for age, gender, and clinical characteristics

We coded studies as follows: y (baseline assessments for age, gender, and clinical characteristics), p (at least one type of assessment), and NR (not reported or unclear). Most studies (121/177 = 68.4%) provided baseline assessments of age, gender, and clinical characteristics. Forty-three studies (24.3%) provided a partial assessment, and 13 studies (7.3%) reported no relevant data.

Baseline comparability of intervention and control groups

We coded studies as follows: y (comparability demonstrated, low risk of bias), p (comparability partially demonstrated, risk of bias unclear), and NR (comparability not demonstrated or unclear, risk of bias unclear or high, depending on whether groups were clearly non-comparable). Studies could demonstrate comparability by reporting data for each group, by reporting the outcome of statistical tests (e.g. P values), or both. One hundred and sixteen studies (65.5%) demonstrated that the groups were comparable at baseline; 27 studies (15.3%) demonstrated partial comparability; and 34 studies (19.2%) did not clearly demonstrate comparability between the groups. No study found that groups were non-comparable (high risk of bias).

Data extraction method for the review

To minimise errors and reduce potential biases being introduced by review authors, the recommended approach is that data extraction should be undertaken independently by at least two people, preferably from complementary disciplines (Section 7.6.2, Higgins 2011). However, in this review, one reviewer (AM) extracted the data, and another reviewer (HH) checked these data.

Effects of interventions

Primary outcome measures

The review analyses 19 comparisons. Of these, 8 are topical treatment versus placebo analyses, and 11 are head-to-head analyses of a topical treatment against a vitamin D analogue (i.e. 1 commonly used class of treatments). Some analyses are a 'catch all' category; for example, analysis 6 includes 'Other treatment versus placebo', which covers 26 treatments for body psoriasis for which there is less research evidence (both in terms of numbers of studies and numbers of participants contributing data). Similarly, analysis 15 incorporates 12 head-to-head comparisons of vitamin D analogues for body psoriasis that are not easily classified under the other head-to-head comparisons. Scalp trials (comparisons 18 and 19) and trials of inverse psoriasis (comparisons 16 and 17) are analysed separately from the trials of body psoriasis.

Table 4 summarises the 19 analyses. Table 2 gives details of the outcome measures considered. The number of participants and number of studies are adjusted manually from those reported in the Tables and Figures to allow for within-patient studies, studies contributing more than once to a single analysis, and studies contributing to multiple analyses. Therefore, numbers of participants and studies reported sometimes differ from the numbers estimated by RevMan.

For each of the 19 analyses, we analysed data on 5 effectiveness outcome measures, where available. The fifth measure is a 'combined end point' that uses data from the four primary outcome measures.

(a) Investigator's Assessment of Overall Global Improvement (IAGI)/Investigator's Global Assessment of Disease Severity (IGA)

Analysis 1: Vitamin D analogues versus placebo

This comparison included eight vitamin D analogues for body psoriasis (see Analysis 1.1 and Table 5). Twenty trials with 3771 participants reported IAGI data on 7 of these treatments. Thirteen trials were between-patient design, and 7 were within-patient studies. Treatment duration ranged from 4 weeks to 12 weeks. The pooled SMD across all treatments was -0.95 (95% CI -1.17 to - 0.74; I² statistic = 89.0%), but there was considerable variation between treatments, so we removed pooling across subgroups. Six treatments were significantly more effective than placebo, with the effect size ranging from -0.67 (becocalcidiol twice daily) to -1.66 (paricalcitol once daily). There was considerable between-study variation in the IAGI SMD for calcitriol. The pooled effect was -1.03 (95% CI -1.71 to -0.36), but this ranged from -0.26 (95% CI -0.99 to 0.47) for Langner 2001 (P) to -3.11 (95% CI -3.57 to -2.66) for Perez 1996. The magnitude of the IAGI SMD for the Perez study was the highest across all comparisons and treatments. For the 'combined end point' of this analysis, we explored the impact of removing this trial from the pooled findings using sensitivity analysis. The presence of considerable heterogeneity within this subgroup means that the estimated average benefit should be interpreted with caution (Higgins 2011).

Analysis 2: Corticosteroid (potent) versus placebo

This comparison included 10 potent corticosteroids for body psoriasis (see Analysis 2.1 and Table 6), although no effectiveness data were available for budesonide. Nine studies with 1867 participants reported IAGI data on 6 of these 10 treatments. Eight trials were between-patient design, and one was a within-patient study (Stein 2001). Treatment duration ranged from 3 to 12 weeks. The SMD across all 6 treatments for IAGI was -1.00 (95% CI -1.18 to -0.82; I² statistic = 57.6%). All six treatments performed statistically significantly better than placebo.

Analysis 3: Corticosteroid (very potent) versus placebo

This comparison included three very potent corticosteroids for treatment of psoriasis of the body (see Analysis 3.1 and Table 7). Five studies with 515 participants reported IAGI data on 2 of the 3 treatments. There were four between-patient trials and 1 within-patient study (Beutner 2006). Treatment duration ranged from two to four weeks. The IAGI SMD across both treatments was -1.87 (95% CI -2.38 to -1.36; I² statistic = 78.7%). Both clobetasol propionate and halobetasol performed statistically significantly better than placebo. In Analysis 18.1, we present placebo-controlled scalp trials of very potent corticosteroids.

Analysis 4: Dithranol versus placebo

Our review did not identify any study comparing dithranol against placebo and which reported IAGI data.

Analysis 5: Vitamin D combination products versus placebo

This comparison included treatment with combined calcipotriol and betamethasone dipropionate used either once or twice daily on the body (see Analysis 5.1 and Table 8). Five parallel-group studies with 2058 participants contributed data on both dosing options. Treatment duration ranged from four to eight weeks. The IAGI SMD across treatments was -1.44 (95% CI -1.76 to -1.12; I² statistic = 89.4%), with twice-daily combination treatment (SMD -1.90; 95% CI -2.09 to -1.71) achieving a significantly larger effect than once-daily treatment (SMD -1.21; 95% CI -1.50 to -0.91). However, the difference between once- and twice-daily dosing was not statistically significant when benefit was assessed using the PASI (see Analysis 5.3). In Analysis 18.1, we report placebocontrolled trials of combination vitamin D/steroid treatments for scalp psoriasis.

Analysis 6: Other treatment versus placebo

This comparison comprised all other treatments for psoriasis of the body not included in the first five analyses; therefore, we removed pooling. None of the studies assessed the same treatment, which means that findings should be interpreted with caution.

In total, we included 26 treatments in this analysis (see Analysis 6.1 and Table 9). Eight studies with 364 participants reported IAGI data on 8 of these 26 treatments. Four trials were betweenpatient design, and four were within-patient studies. Treatment duration ranged from 3 to 12 weeks.

Four treatments performed statistically significantly better than placebo: anti-IL-8 monoclonal antibody cream; betamethasone 17-valerate 21-acetate plus tretinoin plus salicylic acid, indigo naturalise 1.4% ointment, and methotrexate gel. The effect size for the IAGI ranged from -0.56 (Sutton 2001; methotrexate gel) to - 2.14 (Lin 2008; indigo naturalise 1.4% ointment).

In four treatments, the difference relative to placebo was not statistically significant: hexafluoro-1,25-dihydroxyvitamin D3, kukui nut oil, oleum horwathiensis, and platelet aggregation activating factor (PAF). No treatment was statistically significantly less effective than placebo.

Analysis 7: Vitamin D analogues versus corticosteroid (potent)

This comparison included eight vitamin D analogue-potent corticosteroid contrasts for body psoriasis (see Table 10). Eight studies with 2655 participants reported IAGI data for 6 of the 8 intervention-comparator contrasts (see Analysis 7.1). Seven trials were between-patient design, and one was a within-patient study (Medansky 1996). Treatment duration ranged from three to eight weeks. Overall, there was no statistically significant difference between vitamin D analogues and potent corticosteroids: The SMD across all 6 treatments for IAGI was 0.17 (95% CI -0.04 to 0.37; I^2 statistic = 83.4%). In light of the high level of heterogeneity and inconsistency across treatments (Higgins 2011). We removed pooling. Vitamin D analogues performed statistically significantly better than one potent corticosteroid. This finding came from a single between-patient study in which 99 participants contributed data (Bruce 1994). The SMD for calcipotriol against fluocinonide 0.05% ointment was -0.58 (95% CI -0.99 to -0.18; I2 statistic =

NA). Calcipotriol was statistically significantly less effective than both diflorasone diacetate 0.05% ointment (SMD 0.27; 95% CI 0.02 to 0.52) and betamethasone dipropionate (SMD 0.43; 95% CI 0.28 to 0.58; I² statistic = 50.3%). We found no statistically significant difference between calcipotriol and betamethasone valerate, calcitriol and betamethasone dipropionate, or calcitriol and betamethasone valerate.

Analysis 8: Vitamin D analogues versus corticosteroid (very potent)

This comparison included one vitamin D analogue, calcipotriol ointment, versus a very potent corticosteroid contrast, clobetasol propionate foam, for psoriasis of the body (see Table 11 and Analysis 8.1). One study with 42 participants reported IAGI data. Koo 2006 was a between-patient study with a treatment duration of two weeks, which found no significant difference between the treatments (SMD 0.19; 95% CI -0.42 to 0.80).

Analysis 9: Vitamin D combined with corticosteroid versus corticosteroid

This comparison considered vitamin D analogues-steroid combination against potent or very potent corticosteroids for body psoriasis (see Analysis 9.1 and Table 12). The comparison included three contrasts: calcipotriol plus betamethasone dipropionate versus betamethasone dipropionate, calcipotriol plus betamethasone dipropionate versus clobetasol propionate, and calcipotriol plus clobetasol propionate versus clobetasol propionate.

Four between-patient trials reported IAGI data for 1991 participants on 2 of the 3 intervention-comparator contrasts. Treatment duration ranged between two and eight weeks. As these treatment comparisons were very different, we only pooled subtotals. In all but one trial (Fleming 2010 (H)), combination treatment was significantly more effective than corticosteroid alone. Three of the four trials compared calcipotriol/betamethasone dipropionate against betamethasone dipropionate (SMD -0.40; 95% CI -0.52 to -0.27; I² statistic = 41.8%), and one trial compared combined treatment with calcipotriol and clobetasol against clobetasol alone (SMD -0.69; 95% CI -1.22 to -0.15).

Analysis 10: Vitamin D alone or in combination versus dithranol

This comparison considered vitamin D analogues against dithranol (Analysis 10.1 and Table 13). We identified three intervention-comparator contrasts: calcipotriol versus dithranol, calcitriol versus dithranol, and tacalcitol versus dithranol. Five between-patient trials reported IAGI data for 1108 participants on 2 of these 3 intervention-comparator contrasts. Treatment duration ranged from 8 weeks to 12 weeks. There was some variation in the dithranol regimens employed by trials and in the baseline severity of trial participants. These factors may help explain the high level of heterogeneity found in the pooled results.

The SMD for the IAGI was -0.24 (95% CI -0.72 to 0.25; I² statistic = 93.0%). The presence of considerable heterogeneity means that the estimated average benefit should be treated with caution and pooling was therefore removed (Higgins 2011). Data from four trials contributed to the SMD for the calcipotriol versus dithranol: -0.43 (95% CI -0.85 to -0.01; I² statistic = 89.3%), indicating that calcipotriol was statistically significantly more effective than dithranol. Three of these four trials found a significant difference in favour of calcipotriol (Berth Jones 1992b; Christensen 1999; Wall 1998), but the trial by Van de Kerkhof 2006 found outpatient treatment with short contact dithranol to be significantly more effective than calcipotriol alone.

Data from one trial contributed to the SMD for the calcitriol versus dithranol: $0.51 (95\% \text{ CI } 0.13 \text{ to } 0.88; \text{I}^2 \text{ statistic} = \text{NA})$, indicating that dithranol was statistically significantly more effective than calcitriol.

Analysis 11: Vitamin D alone or in combination versus other vitamin D analogue

Our review identified three intervention-comparator contrasts in this comparison: calcipotriol versus calcitriol, calcipotriol versus tacalcitol, and calcipotriol versus maxacalcitol for body psoriasis (see Analysis 11.1 and Table 14). Three trials involving 498 participants contributed IAGI data for all 3 intervention-comparator contrasts (one trial for each contrast). Two trials were betweenpatient, and one was within-patient in design (Barker 1999 (H)). Treatment duration ranged from 8 to 12 weeks. The SMD for the IAGI was -0.06 (95% CI -0.51 to 0.38; I² statistic = 82.2%). The presence of substantial heterogeneity reflects differences in the findings from the two intervention-comparator contrasts underlying this statistic. We found a statistically significant difference in favour of calcipotriol in the analysis against tacalcitol (SMD -0.47; 95% CI -0.73 to -0.21), but there was no significant difference relative to calcitriol (SMD 0.00; 95% CI -0.25 to 0.25) or between calcipotriol and maxacalcitol (SMD 0.43; 95% CI -0.12 to 0.98). In light of the inconsistency of results across treatments, we only pooled subtotals (Higgins 2011).

Analysis 12: Vitamin D alone or in combination versus vitamin D + corticosteroid

Our review identified 12 intervention-comparator contrasts for body psoriasis, involving 3 vitamin D analogues, 2 combination products, and 7 different corticosteroids (see Analysis 12.1 and Table 15). Eleven parallel-group trials involving 4791 participants contributed IAGI data for 9 of these 12 intervention-comparator contrasts. Treatment duration ranged from two to eight weeks. There were no IAGI data for three contrasts: calcipotriol versus calcipotriol plus diflucortolone valerate, calcipotriol versus calcipotriol plus fluocinonide acetonide, and calcitriol versus calcitriol plus diflucortolone valerate.

Overall, vitamin D alone appeared to be less effective than vitamin D plus corticosteroid: The SMD for the IAGI was 0.48 (95% CI 0.32 to 0.65; I² statistic = 86.9%). This finding applied to all but two of the intervention-comparator contrasts: There was no statistically significant difference between twice-daily calcipotriol and a regimen of calcipotriol (morning) plus betamethasone valerate (night time) (SMD 0.27; 95% CI -0.19 to 0.74) (Kragballe 1998b), or between once-daily calcipotriol and once-daily treatment with a combined product containing calcipotriol and hydrocortisone (SMD 0.14; 95% CI -0.06 to 0.33) (Ortonne 2010). The study by Ortonne 2010 also reported findings separately for the face (see Analysis 17.1).

In light of the observed heterogeneity between the interventioncomparator contrasts, we only pooled subtotals (Higgins 2011).

Analysis 13: Vitamin D alone or in combination versus other treatments: complex regimens

This comparison summarises findings on complex regimens for body psoriasis, defined here as treatment sequences that do not consist of a simple head-to-head comparison between two active treatments (see Analysis 13.1 and Table 16). We identified 12 intervention-comparator contrasts, and IAGI data were available for 10 of these. Data from 2755 participants contributed to the IAGI analysis, based on findings from seven trials. Six of the trials were between-patient (parallel-group) in design, and one was a within-patient study (Austad 1998). Trial duration varied between 6 and 12 weeks. As the interventions and comparators were highly variable and because two trials each contributed three pair-wise contrasts, we did not pool the data.

Six intervention-comparator contrasts for which IAGI data were available found a significant difference between regimens. A twoweek regimen with clobetasol propionate followed by four weeks' treatment with calcipotriol was more effective than six weeks of monotherapy with calcipotriol (SMD 0.60; 95% CI 0.18 to 1.02). All treatments were applied twice-daily in this within-patient study (Austad 1998).

The remaining five more effective regimens all involved once-daily treatment with a combined product containing calcipotriol and betamethasone dipropionate, and the same study (White 2006 (P)) reported three contrasts. This study involved an initial 'treatment' phase consisting of 4 weeks' treatment with the combined product, followed by an 8-week maintenance phase. In one maintenance phase, participants received placebo ointment for 8 weeks. This was significantly less effective than maintenance with once-daily calcipotriol (SMD 0.27; 95% CI 0.12 to 0.41), and it was also less effective than maintenance with once-daily calcipotriol on weekdays and the combined product at the weekend (SMD 0.51; 95% CI 0.37 to 0.66). When these two comparator regimens were compared directly (White 2006 (H)), the alternating

regimen using weekday/weekend treatments during the maintenance phase was significantly more effective than once-daily calcipotriol for maintenance (SMD 0.26; 95% CI 0.11 to 0.40). Kragballe 2004 compared 3 12-week treatment regimens, 2 of which were 'complex'. When we compared these complex regimens, eight weeks' once-daily combination treatment with calcipotriol and betamethasone dipropionate, followed by four weeks of once-daily calcipotriol was significantly less effective than a regimen consisting of four weeks' once-daily combination treatment, followed by eight weeks of once-daily calcipotriol on weekdays and the combined product at weekends (SMD 0.24; 95% CI 0.08 to 0.40). A separate trial compared 8 weeks with tacalcitol to combination ointment for 4 weeks, followed by calcipotriol for 4 weeks (Ortonne 2004). Participants applied all treatments once daily (at night time). Monotherapy with tacalcitol was significantly less effective than the complex regimen (SMD 0.54; 95% CI 0.36 to 0.72).

In 4 of the 10 intervention-comparator contrasts for which IAGI data were available, there was no significant difference in effect. The study by Kragballe 2004 (see paragraph above) found no difference between twice-daily calcipotriol and the two comparator complex regimens. One regimen consisted of eight weeks' oncedaily treatment with a combined product containing calcipotriol and betamethasone dipropionate, followed by four weeks of oncedaily calcipotriol. Relative to calcipotriol monotherapy, the SMD for the IAGI was -0.12 (95% CI -0.29 to 0.04). When compared with a regimen consisting of four weeks' once-daily treatment with a combined product containing calcipotriol and betamethasone dipropionate, followed by eight weeks of once-daily calcipotriol on weekdays and the combined product at weekends, twice-daily calcipotriol appeared to be slightly less effective, although the difference was not statistically significant (SMD 0.13; 95% CI -0.04 to 0.29).

Yang 2009 compared a six-week course of twice-daily calcipotriol with a complex routine using halometasone and calcipotriol. For two weeks participants applied halometasone in the morning and calcipotriol at night; over the next two weeks, halometasone was applied twice daily at weekends and calcipotriol twice daily during the week. For the final two weeks, treatment reverted to twice-daily calcipotriol. Although there was a trend in favour of the complex regimen (SMD 0.41; 95% CI -0.05 to 0.86), the difference was not statistically significant.

Lahfa 2003 compared two different vitamin D products combined with a very potent corticosteroid. In the initial phase, participants applied dual therapy (clobetasol propionate in the mornings and the vitamin D analogue (calcipotriol or calcitriol) at night) until achieving clearance or marked improvement or until 4 weeks had elapsed. Participants then switched to a monotherapy maintenance phase with the vitamin D product for the remainder of the 12-week study period. The IAGI results showed the 2 treatment regimens to be equivalent: -0.19 (95% CI -0.54 to 0.16).

Analysis 14: Vitamin D alone or in combination versus other treatment: long-term studies (> 24 weeks)

This comparison included active-controlled studies of psoriasis of the body that were at least 24 weeks in duration (see Analysis 14.1 and Table 17). One longer-term placebo-controlled trial of the body (Katz 1991a) and two long-term scalp trials (Luger 2008; Poulin 2010) did not meet the inclusion criteria for this comparison, and they are evaluated elsewhere (Analysis 2, Analysis 19, and Analysis 18, respectively).

One trial was included in this comparison; Kragballe 2006 was a between-patient trial that compared three 52-week regimens with all treatments used once daily:

• first, combination treatment with calcipotriol and betamethasone dipropionate;

• second, alternating treatment with combination therapy for 4 weeks, then calcipotriol for 4 weeks; and

• third, combination therapy for 4 weeks, then calcipotriol for 48 weeks.

In total, 297 participants contributed IGA data to the analysis. Data were unsuitable for pooling because the same participants contributed to more than one analysis.

According to the investigators' assessment, there was no significant difference between the three long-term regimens. One year's combination therapy was not significantly better than either the alternating regimen (SMD -0.09; 95% CI -0.36 to 0.18) or the regimen of treatment with 4 weeks of combination therapy followed by 48 weeks of calcipotriol (SMD -0.18; 95% CI -0.47 to 0.10). In both these comparisons, combination therapy achieved a larger absolute benefit, but the difference was not statistically significant. When alternating therapy was compared with the regimen of 4 weeks' combination therapy followed by 48 weeks of calcipotriol, the SMD for the IAGI also indicated the two regimens were not statistically significantly different: -0.09 (95% CI -0.37 to 0.19).

Analysis 15: Vitamin D analogues versus other treatment

This comparison incorporated all other vitamin D head-to-head comparisons of treatments for psoriasis of the body (excluding inverse psoriasis) that had not already been included (see Analysis 15.1 and Table 18). We included 12 intervention-comparator contrasts, with IAGI data available for 6 of these contrasts. Eight between-patient trials and 2 within-patient trials provided data for 1386 participants. Trial duration ranged between 4 and 12 weeks. In light of the pharmacological diversity amongst the comparators, we only pooled data within subgroups.

According to the investigator's global assessment, twice-daily calcipotriol was significantly more effective than coal tar polytherapy (SMD -0.59; 95% CI -0.87 to -0.31; I² statistic = 0%). Oncedaily vitamin D was significantly less effective than a twice-daily application (SMD -0.24; 95% CI -0.38 to -0.09; I² statistic = 0%). This contrast included a comparison of once- versus twicedaily dosing of calcipotriol (SMD -0.27; 95% CI -0.48 to -0.06) and a dosing comparison combination treatment with calcipotriol and betamethasone dipropionate (SMD -0.20; 95% CI -0.41 to 0.00).

In the remaining four intervention-comparator contrasts, no significant difference was found between twice-daily calcipotriol and the comparators. This finding held for the comparison against coal tar monotherapy (SMD -0.53; 95% CI -1.74 to 0.68; I² statistic = 91.1%), against betamethasone dipropionate ointment and salicylic acid (SMD -0.06; 95% CI -0.33 to 0.22; I² statistic = NA), against tacrolimus ointment (SMD -0.22; 95% CI -0.60 to 0.16; I² statistic = NA), and against vitamin B12 cream (SMD -0.55; 95% CI -1.33 to 0.24; I² statistic = NA).

Two of the three trials comparing calcipotriol with coal tar monotherapy found a significant difference in favour of calcipotriol; the other trial found a significant difference in favour of coal tar (Alora-Palli 2010). This apparent contradiction may reflect different formulations of coal tar, treatment durations, or different baseline disease severity.

Analysis 16: Flexural/facial psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of topical treatments for inverse or facial psoriasis (see Analysis 16.1 and Table 19). Evidence on four treatments was found in this comparison: the potent steroid betamethasone valerate; the vitamin D analogue calcipotriol; and two topical calcineurin inhibitors, pimecrolimus and tacrolimus. We only found one trial that evaluated tacrolimus ointment (Lebwohl 2004), but the study did not report any effectiveness data suitable for this review. However, the study did contribute data on adverse events and withdrawal rates. IAGI outcome data were available for one of the four topical treatments for inverse psoriasis. One 8-week between-patient study with 47 participants (Gribetz 2004) reported placebo-controlled data on pimecrolimus 1% cream, which demonstrated a statistically significant difference in favour of twice-daily pimecrolimus (SMD -1.07; 95% CI -1.69 to -0.45).

Analysis 17: Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for inverse psoriasis, which compared vitamin D with an active control (see Analysis 17.1 and Table 20). We identified five intervention-comparator contrasts. Four treatments were compared with calcipotriol: once-daily betamethasone valerate, combined treatment with calcipotriol and hydrocortisone, calcitriol, and pimecrolimus. Calcitriol was compared with tacrolimus.

Two between-patient studies contributed IAGI data from 457 study participants on 2 of the 5 intervention-comparator contrasts. Trial duration ranged from six to eight weeks. When applied once daily to inverse psoriasis, calcipotriol was significantly less effective than combined treatment with calcipotriol and hydrocortisone (SMD 0.30; 95% CI 0.11 to 0.50) (Ortonne 2010). However, there was no significant difference in effect between twice daily treatment with calcitriol and tacrolimus (SMD 0.42; 95% CI - 0.15 to 0.98) (Liao 2007).

Analysis 18: Scalp psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of treatments for scalp psoriasis (see Analysis 18.1 and Table 21). We included evidence on 11 treatments in this comparison, with IAGI data available for 9 treatments. Ten trials, 9 of which were between-patient studies, contributed data from 2472 participants. One study was a within-patient trial (Lepaw 1978). Trial duration ranged between two and eight weeks. IAGI data were not available for betamethasone valerate or ciclopirox olamine shampoo.

Eight treatments for scalp psoriasis that were assessed using the IAGI scale were significantly more effective than placebo. The least effective treatment was calcipotriol (SMD -0.72; 95% CI -1.28 to -0.16; I² statistic = 69.2%), and the most effective treatment was clobetasol propionate (SMD -1.73; 95% CI -1.99 to -1.48; I² statistic = 14.3%). Combination treatment with calcipotriol and betamethasone dipropionate was also effective (SMD -0.97; 95% CI -1.61 to -0.32; I² statistic = 90.2%). Data from the three very potent corticosteroids were pooled (amcinonide, clobetasol propionate, halcinonide). The SMD across these treatments was -1.57 (95% CI -1.85 to -1.28; I² statistic = 49.7%). Only salicylic acid was not significantly more effective than placebo (SMD -0.86; 95% CI -1.79 to 0.06).

Analysis 19: Scalp psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for scalp psoriasis in which one of the interventions was a vitamin D product (used either as monotherapy or in combination with another product) (see Analysis 19.1 and Table 22). We identified six intervention-comparator contrasts, and IAGI data were available for five of these contrasts. All studies were parallel-group in design (between-patient). Ten studies contributed IAGI data from 5175 participants, and trial duration ranged from 4 to 52 weeks. Vitamin D for scalp psoriasis was significantly less effective than potent steroids, either alone or in combination with vitamin D. Specifically, calcipotriol was less effective than three comparator treatments: betamethasone dipropionate (SMD 0.48; 95% CI 0.32 to 0.64; I² statistic = 60.4%), betamethasone valerate (SMD 0.37; 95% CI 0.20 to 0.55; I² statistic = 0%), and combination treatment with calcipotriol and betamethasone dipropionate (SMD 0.64; 95% CI 0.44 to 0.84; I² statistic = 82.3%). Combination treatment (calcipotriol/betamethasone dipropionate) was significantly more effective than betamethasone dipropionate alone (SMD -0.18; 95% CI -0.26 to -0.10; I² statistic = 0%). The efficacy of calcipotriol and coal tar polytherapy was similar: SMD - 0.24 (95% CI -0.73 to 0.25; I² statistic = 91.1%).

(b) Total Severity Scores (TSS)

Analysis 1: Vitamin D analogues versus placebo

Our review included eight vitamin D analogues for body psoriasis in this comparison (see Analysis 1.2 and Table 5). Nineteen studies reported TSS data with 2647 participants contributing data. Nine trials were between-patient design, and 10 were within-patient studies. Treatment duration ranged from 4 weeks to 12 weeks. The average effect size across all 8 treatments for TSS was -1.04; (95% CI -1.33 to -0.74; I² statistic = 93.0%), but two of these treatments were not significantly more effective than placebo. The high level of heterogeneity means that the estimated average benefit should be treated with caution; there was considerable variation between effective treatments, with the TSS ranging from -0.46 (becocalcidiol twice daily) to -2.15 (paricalcitol once daily). Therefore, we removed pooling across subgroups (Higgins 2011). Four studies contributed data for the SMD for the TSS of calcitriol (SMD -1.22; 95% CI -2.38 to -0.07), but there was considerable heterogeneity within the subgroup (I² statistic = 98.3%). One of the studies (Van de Kerkhof 1989) found no statistically significant difference between calcitriol and placebo (SMD -0.06; 95% CI -0.94 to 0.81), whereas the study by Perez 1996 found a large and statistically significant difference (-4.03; 95% CI -4.56 to -3.50). We considered this finding in more detail in the 'combined end point' section (e) below.

Analysis 2: Corticosteroid (potent) versus placebo

Our review included 10 potent corticosteroids for body psoriasis in this comparison (see Analysis 2.2 and Table 6). Seven studies reported TSS data contributed by 553 participants on 8 of these 10 treatments. Six trials were between-patient studies, and there was one within-patient design (Ormerod 1997). Treatment duration ranged from 2 to 12 weeks. The average effect size across all 8 treatments for TSS was -0.77 (95% CI -1.01 to -0.52; I² statistic = 46.7%). We found all treatments, except for diflorasone diacetate, to be statistically significantly superior to placebo: SMD -0.32; 95% CI -0.73 to 0.09).

Analysis 3: Corticosteroid (very potent) versus placebo

Of the three very potent corticosteroids for body psoriasis included in this comparison, TSS data were available only for clobetasol propionate (see Analysis 3.2 and Table 7). Three studies, all of which were between-patient trials, reported TSS data for 545 participants. Treatment duration ranged from two to four weeks. The SMD for the TSS was -1.35 (95% CI -1.80 to -0.89; I² statistic = 75.3%). In Analysis 18.2, we reported TSS data on the use of very potent steroids on the scalp.

Analysis 4: Dithranol versus placebo

This comparison considered dithranol against placebo (see Analysis 4.2 and Table 23). Three within-patient trials reported TSS data for 47 participants. Treatment duration ranged from three to eight weeks. The SMD for the TSS was -1.06 (95% CI - 1.66 to -0.46; I² statistic = 37.4%).

Analysis 5: Vitamin D combination products versus placebo

Our review did not identify any trial reporting TSS data for this comparison.

Analysis 6: Other treatment versus placebo

This comparison comprised all other treatments for body psoriasis that were not included in the first five comparisons. None of the studies assessed the same treatment, which means that findings should be interpreted with caution.

In total, we included 26 treatments in this analysis (see Analysis 6.2 and Table 9). Seventeen studies with 907 participants reported TSS data on 17 of these 26 treatments. Five trials were betweenpatient design, and 12 were within-patient studies. Treatment duration ranged from 3 to 12 weeks.

Ten treatments performed significantly better than placebo: anti-IL-8 monoclonal antibody cream, calcipotriene 0.005% ointment + nicotinamide, fish oil plus occlusion, hexafluoro-1,25-dihydroxyvitamin D3, indigo naturalise 1.4% ointment, methotrexate gel, mycophenolic acid ointment, oleum horwathiensis, PTH (1-34) in Novasome cream®, and tazarotene. The effect size for the TSS ranged from -0.48 (Levine 2010 (P); calcipotriene 0.005% ointment + nicotinamide) to -2.31 (Holick 2003; PTH (1-34) in Novasome cream®).

In seven treatments, the difference relative to placebo was not statistically significant: kukui nut oil, NG-monomethyl-L-arginine (L-NMMA) cream, nicotinamide 1.4%, polymyxin B cream, topical sirolimus, topical tacrolimus, and tar.

Analysis 7: Vitamin D analogues versus corticosteroid (potent)

There were eight vitamin D analogue-potent corticosteroid comparisons for body psoriasis (see Analysis 7.2 and Table 10). Six studies with 891 participants reported TSS data for 5 of the 8 intervention-comparator contrasts. Two trials were between-patient design, and four were within-patient studies. Treatment duration ranged from three to six weeks. The SMD across all 5 treatments for TSS indicated that there was no significant difference between the vitamin D derivatives and potent corticosteroid: SMD 0.11 (95% CI -0.22 to 0.44; I² statistic = 86.7%). However, there was considerable variation between the individual contrasts underlying the pooled effect. In light of this heterogeneity, we removed pooling (Higgins 2011).

In two of the five vitamin D-potent corticosteroid comparisons, the vitamin D analogue performed statistically significantly better than the potent corticosteroid: The SMD for calcipotriol against fluocinonide 0.05% ointment was -0.50 (95% CI -0.92 to -0.07; I² statistic = NA) (Bruce 1994). Similarly, the comparison of calcipotriol against betamethasone valerate showed a significant difference in favour of calcipotriol (SMD -0.26; 95% CI -0.41 to -0.11), a finding also based on a single study (Kragballe 1991a). In three comparisons, the vitamin D analogue was statistically significantly less effective than the potent corticosteroid: calcipotriol versus difforasone diacetate (SMD 0.40; 95% CI 0.15 to 0.65), calcitriol versus betamethasone dipropionate (SMD 0.27; 95% CI 0.02 to 0.51), and tacalcitol versus betamethasone valerate (SMD 0.41; 95% CI 0.09 to 0.74).

Analysis 8: Vitamin D analogues versus corticosteroid (very potent)

Our review did not identify any study of psoriasis of the body that compared vitamin D analogues against very potent corticosteroids and that reported TSS data.

Analysis 9: Vitamin D combined with corticosteroid versus corticosteroid

This comparison considered vitamin D analogues-steroid combination against potent or very potent corticosteroid for body psoriasis (see Analysis 9.2 and Table 12). One four-week betweenpatient trial reported TSS data for 122 participants with moderate to severe psoriasis (Menter 2009). The combined treatment with calcipotriol and betamethasone dipropionate was found to be significantly less effective than clobetasol propionate spray alone (0.45; 95% CI 0.09 to 0.81).

Analysis 10: Vitamin D alone or in combination versus dithranol

This comparison considered vitamin D analogues against dithranol (see Analysis 10.2 and Table 13). We identified three intervention-comparator contrasts: calcipotriol versus dithranol, calcitriol versus dithranol, and tacalcitol versus dithranol. Three betweenpatient trials and 1 within-patient trial (Grattan 1997 (H)) reported TSS data for 386 participants. Treatment duration ranged from four weeks to eight weeks. There was some variation in the dithranol regimens employed by trials and in the baseline severity of trial participants. These factors may help explain the substantial heterogeneity found in the pooled results (Higgins 2011).

The SMD for the TSS was -0.27 (95% CI -0.73 to 0.20; I² statistic = 80.6%). Data from two trials contributed to the SMD for the calcipotriol versus dithranol: -0.54 (95% CI -1.16 to 0.08; I² statistic

= 71.2%) (Christensen 1999; Grattan 1997 (H)). Data from one trial contributed to the SMD for the calcitriol versus dithranol: 0.13 (95% CI -0.24 to 0.50; I² statistic = NA) (Hutchinson 2000). Data from one trial contributed to the SMD for the tacalcitol versus dithranol: -0.18 (95% CI -0.60 to 0.25; I² statistic = NA) (Farkas 1999). Therefore, neither the summary statistic for the TSS nor the pooled data for individual intervention-comparator contrasts provided evidence of a statistically significant advantage of a vitamin D analogue over dithranol or vice versa.

Analysis 11: Vitamin D alone or in combination versus other vitamin D analogue

Our review identified three intervention-comparator contrasts for body psoriasis in this comparison: calcipotriol versus calcitriol, calcipotriol versus tacalcitol, and calcipotriol versus maxacalcitol (see Analysis 11.2 and Table 14). Three trials involving 563 participants contributed TSS data for all 3 of these interventioncomparator contrasts. Two trials were between-patient, and one was within-patient in design (Barker 1999 (H)). Treatment duration ranged from 8 to 12 weeks. The SMD for the TSS indicated that there was a statistically significant difference in favour of calcipotriol: SMD -0.31; 95% CI -0.55 to -0.06; I² statistic = 46.9%. When assessed by the TSS, calcipotriol was statistically significantly more effective than calcitriol (SMD -0.32; 95% CI -0.57 to -0.07). This finding contrasts with the IAGI assessment from the same study, which found no significant difference (see Ji 2008; Analysis 11.1). Calcipotriol was also significantly more effective than tacalcitol (SMD -0.45; 95% CI -0.68 to -0.22) and similar in efficacy relative to maxacalcitol (SMD 0.13; 95% CI -0.41 to 0.68).

Analysis 12: Vitamin D alone or in combination versus vitamin D + corticosteroid

Our review identified 12 intervention-comparator contrasts for body psoriasis, involving 3 vitamin D analogues, 2 combination products, and 7 different corticosteroids (see Analysis 12.1 and Table 15). One 4-week parallel-group trial contributed TSS data from 301 participants for 1 of these 12 intervention-comparator contrasts (Huang 2009). Twice-daily calcipotriol was found to be statistically significantly less effective than once-daily treatment with a combined product containing calcipotriol and betamethasone dipropionate (SMD 0.25; 95% CI 0.03 to 0.48).

Analysis 13: Vitamin D alone or in combination versus other treatments: complex regimens

This comparison summarises findings on complex regimens for body psoriasis, defined here as treatment sequences that do not consist of a simple head-to-head comparison between two active treatments (see Analysis 13.2 and Table 16). We identified 12 intervention-comparator contrasts and TSS data were available for one of these. Data from 1 6-week within-patient trial with 46 participants (Austad 1998) contributed to the TSS analysis. Austad 1998 compared six weeks of twice-daily calcipotriol with a regimen of two weeks' treatment with clobetasol propionate followed by four weeks with calcipotriol. The complex regimen was significantly more effective than monotherapy with calcipotriol (SMD 0.63; 95% CI 0.21 to 1.05).

Analysis 14: Vitamin D alone or in combination versus other treatment: long-term studies (> 24 weeks)

We did not identify any relevant study that provided TSS data for this comparison.

Analysis 15: Vitamin D analogues versus other treatment

This comparison incorporated all other vitamin D head-to-head comparisons of treatments for psoriasis of the body (excluding inverse psoriasis) that had not already been included (see Analysis 15.2 and Table 18). We included 12 intervention-comparator contrasts, with TSS data available for 6 of these contrasts. Five between-patient trials and 2 within-patient trials provided data from 898 participants. Trial duration ranged between 6 and 12 weeks. In light of the pharmacological diversity of the comparators, we only pooled data within subgroups.

According to the TSS assessment, twice-daily calcipotriol was significantly more effective than coal tar polytherapy (SMD -0.51; 95% CI -0.86 to -0.16; I² statistic = NA). In the remaining five intervention-comparator contrasts, we found no significant difference between twice-daily calcipotriol and the comparators.

Analysis 16: Flexural/facial psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of topical treatments for inverse or facial psoriasis (see Analysis 16.2 and Table 19). Evidence on four treatments was found in this comparison: the potent steroid betamethasone valerate; the vitamin D analogue calcipotriol; and two topical calcineurin inhibitors, pimecrolimus and tacrolimus. We found only one placebo-controlled trial that evaluated tacrolimus ointment (Lebwohl 2004), but the study did not report any effectiveness data suitable for this review. However, the study did contribute data on adverse events and withdrawal rates.

TSS data were available for one of the four topical treatments for inverse psoriasis. One 8-week between-patient study contributed data on pimecrolimus 1% cream from 57 participants (Gribetz 2004), 10 participants more than those with an investigator's assessment (see Analysis 16.1). Findings from the TSS were consistent with those of the IAGI: the SMD for the TSS also found a statistically significant difference in favour of twice-daily pimecrolimus (SMD -1.37; 95% CI -1.95 to -0.79).

Analysis 17: Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for inverse psoriasis, which compared vitamin D with an active control (see Analysis 17.2 and Table 20). We identified five intervention-comparator contrasts. Four treatments were compared with calcipotriol: once-daily betamethasone valerate, combined treatment with calcipotriol and hydrocortisone, calcitriol, and pimecrolimus. Calcitriol was compared with tacrolimus.

Two 6-week studies contributed TSS data from 124 study participants on 2 of the 5 intervention-comparator contrasts. Calcipotriol was significantly less effective than calcitriol (SMD 0.61; 95% CI 0.28 to 0.94) (Ortonne 2003). In this within-patient study, participants applied treatments twice daily to 'sensitive areas', including the face, hairline, retro-auricular, and flexural areas. When they applied treatments to the facial and genitofemoral areas, there was no significant difference in effect between twice-daily treatment with calcitriol and tacrolimus (SMD 0.29; 95% CI -0.27 to 0.85) (Liao 2007). In this between-patient study, participants applied both treatments twice daily.

Analysis 18: Scalp psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of treatments for scalp psoriasis (see Analysis 18.2 and Table 21). We included evidence on 11 treatments in this comparison, with TSS data available for 10 treatments. Twelve between-patient trials contributed data from 2897 participants. Trial duration ranged between two and eight weeks. TSS data were not available for halcinonide (classed as a very potent corticosteroid).

Eight of the 10 treatments for scalp psoriasis that were assessed using the TSS scale were significantly more effective than placebo. The least effective treatment was calcipotriol (SMD -0.44; 95% CI -0.64 to -0.25; I² statistic = 0%), and the most effective were the 2 very potent steroids, amcinonide (SMD -1.58; 95% CI -1.98 to -1.18) and clobetasol propionate (SMD -1.53; 95% CI -1.77 to -1.28; I² statistic = 46.3%). Combination treatment with calcipotriol and betamethasone dipropionate was also effective (SMD -0.92; 95% CI -1.42 to -0.43; I² statistic = 83.2%).

We pooled data from the two potent corticosteroids, betamethasone valerate and betamethasone dipropionate (SMD -1.13; 95% CI -1.44 to -0.81; I² statistic = 52.2%). We also pooled data from the two very potent corticosteroids, amcinonide and clobetasol propionate (SMD -1.55; 95% CI -1.73 to -1.37; I² statistic = 19.6%).

Two treatments were not significantly different to placebo: ciclopirox olamine shampoo (SMD -0.07; 95% CI -0.82 to 0.68) and salicylic acid (SMD -0.57; 95% CI -1.47 to 0.32).

Analysis 19: Scalp psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for scalp psoriasis in which one of the interventions was a vitamin D product (used either as monotherapy or in combination with another product) (see Analysis 19.2 and Table 22).

We identified six intervention-comparator contrasts, and TSS data were available for all six of these contrasts. All studies were parallelgroup in design (between-patient). Eleven studies contributed TSS data from 4877 participants, and trial duration ranged from 4 to 8 weeks.

Based on Total Severity Scores, calcipotriol was significantly less effective than betamethasone dipropionate (SMD 0.45; 95% CI 0.28 to 0.63; I² statistic = 66.3%), clobetasol propionate (SMD 0.37; 95% CI 0.05 to 0.69; I² statistic = NA), and combination treatment with calcipotriol and betamethasone dipropionate (SMD 0.70; 95% CI 0.56 to 0.84; I² statistic = 49.6%). There was no statistically significant difference between calcipotriol and betamethasone valerate (SMD 0.09; 95% CI -0.09 to 0.27; I² statistic = 0%). Combination treatment (calcipotriol/betamethasone dipropionate) was significantly more effective than betamethasone dipropionate alone (SMD -0.19; 95% CI -0.27 to -0.11; I² statistic = 0%). The efficacy of calcipotriol and coal tar polytherapy was not significantly different (SMD -0.30; 95% CI -0.84 to 0.24; I² statistic = 93.1%).

(c) Psoriasis Area and Severity Index (PASI)

Analysis 1: Vitamin D analogues versus placebo

Our review included eight vitamin D analogues for body psoriasis in this comparison (see Analysis 1.3 and Table 5). Nine studies reported PASI data, with 2357 participants contributing data on 2 (calcipotriol and tacalcitol) of these 8 treatments. Eight trials were between-patient design, and one was a within-patient study (Dubertret 1992). Treatment duration ranged from three weeks to eight weeks. The average effect size across the 2 treatments for PASI was SMD -0.58 (95% CI -0.71 to -0.45; I² statistic = 42.3%).

Analysis 2: Corticosteroid (potent) versus placebo

Our review included 10 potent corticosteroids for body psoriasis in this comparison (see Analysis 2.3 and Table 6). Three between-patient studies reported PASI data from 1158 participants on once-daily or twice-daily doses of betamethasone dipropionate (2 of the 10 potent steroid regimens considered in this group). Trial duration ranged between four and eight weeks. The average PASI effect size across both application frequencies was -0.97 (95% CI -1.31 to -0.62; I² statistic = 79.6%).

Topical treatments for chronic plaque psoriasis (Review)

Analysis 3: Corticosteroid (very potent) versus placebo

Our review did not identify any study that compared very potent corticosteroids against placebo and also reported PASI data. This may be because whole-body application of very potent corticosteroids is not recommended, so a whole-body assessment measure like the PASI is inappropriate.

Analysis 4: Dithranol versus placebo

Our review did not identify any study that compared dithranol against placebo and also reported PASI data.

Analysis 5: Vitamin D combination products versus placebo

This comparison included treatment with combined calcipotriol and betamethasone dipropionate used either once or twice daily on the body (see Analysis 5.3 and Table 8). Five parallel-group studies with 2056 participants contributed PASI data on both dosing options. Treatment duration ranged from four to eight weeks. The PASI SMD across treatments was -1.24 (95% CI -1.53 to -0.95; I² statistic = 87.6%). Although twice-daily combination treatment (SMD -1.41; 95% CI -1.86 to -0.97) achieved a larger effect than once-daily treatment (SMD -1.14; 95% CI -1.57 to -0.70), the difference was not statistically significant at the 5% level. This finding contrasts with the assessment of the relative benefit of once- versus twice-daily dosing using the IAGI metric (see Analysis 5.1).

Analysis 6: Other treatment versus placebo

This comparison comprised all other treatments for body psoriasis that were not included in the first five comparisons; therefore, we removed pooling. None of the studies assessed the same treatment, which means that findings should be interpreted with caution. In total, we included 26 treatments in this analysis (see Analysis 6.3 and Table 9). Nine studies with 529 participants reported

PASI data on 9 of these 26 treatments. Eight trials were betweenpatient studies, and one was within-patient in design (Vali 2005). Treatment duration ranged from 2 to 12 weeks.

Six treatments performed statistically significantly better than placebo: aloe vera extract, betamethasone 17-valerate 21-acetate plus tretinoin plus salicylic acid, a herbal skin care product, *Mahonia aquifolium*, methotrexate gel, and theophylline ointment. Effects sizes ranged from -0.54 (95% CI -0.99 to -0.10) for the betamethasone 17-valerate 21 product (Santoianni 2001) to -2.96 (95% CI -4.19 to -1.74) for the herbal skin care treatment (Maier 2004).

In three treatments, we found no statistically significant difference relative to placebo in the PASI assessments. These comprised topical caffeine (Vali 2005), dead sea salts emollient lotion (Cheesbrough 1992), and kukui nut oil (Brown 2005).

Analysis 7: Vitamin D analogues versus corticosteroid (potent)

Our review identified eight vitamin D analogue-potent corticosteroid comparisons for body psoriasis (see Analysis 7.3 and Table 10). Nine studies with 3185 participants reported PASI data for 4 of the 8 intervention-comparator contrasts. Seven trials were between-patient design, and two were within-patient studies. Treatment duration ranged from four to eight weeks. The PASI SMD across all four intervention-comparator contrasts indicated that there was no statistically significant difference between the vitamin D derivatives and potent corticosteroid (SMD 0.12; 95% CI -0.07 to 0.32; I² statistic = 86.2%).

In one vitamin D analogue versus potent corticosteroid comparison, the vitamin D analogue performed statistically significantly better than the potent corticosteroid: four studies contributed data to analysis of calcipotriol versus betamethasone valerate (SMD -0.12; 95% CI -0.22 to -0.02; I² statistic = 0%).

In two intervention-comparator contrasts, the vitamin D analogue was statistically significantly less effective than the potent corticosteroid: Betamethasone dipropionate was more effective than both calcipotriol (SMD 0.36; 95% CI 0.22 to 0.51; I² statistic = 49.6%) and calcitriol (SMD 0.39; 95% CI 0.14 to 0.63; I² statistic = NA).

We found no statistically significant difference between calcipotriol and desoxymetasone.

Analysis 8: Vitamin D analogues versus corticosteroid (very potent)

This comparison considered vitamin D analogues against very potent corticosteroids for treating psoriasis of the body (see Analysis 8.3 and Table 11). We found data on one intervention-comparator contrast: calcipotriol versus clobetasol propionate. One betweenpatient trial reported PASI data for 40 participants. This trial had a treatment duration of six weeks (Landi 1993). The SMD for the PASI indicated that there was no statistically significant difference between the very potent corticosteroid and the vitamin D analogue: SMD -0.32 (95% CI -0.95 to 0.30; I² statistic = NA).

Analysis 9: Vitamin D combined with corticosteroid versus corticosteroid

This comparison considered vitamin D analogue-steroid combination against potent or very potent corticosteroid for body psoriasis (Table 12). Three between-patient trials reported PASI data for 1876 participants (see Analysis 9.3). All 3 trials compared combination treatment with calcipotriol and betamethasone dipropionate with betamethasone dipropionate as monotherapy, but varied in their treatment duration (4 to 8 weeks). The SMD for the PASI was -0.44 (95% CI -0.55 to -0.33; I² statistic = 22.4%), indicating a significantly greater effect for combination treatment.

Analysis 10: Vitamin D alone or in combination versus dithranol

This comparison considered vitamin D analogues against dithranol (see Analysis 10.3 and Table 13). We identified three intervention-comparator contrasts: calcipotriol versus dithranol, calcitriol versus dithranol, and tacalcitol versus dithranol. Five between-patient trials reported PASI data for 796 participants. Treatment duration ranged from 8 to 12 weeks. There was some variation in the dithranol regimens employed by trials and in the baseline severity of trial participants. These factors may explain the considerable level of heterogeneity found in the pooled results (Higgins 2011). The SMD for the PASI was 0.36 (95% CI -0.33 to 1.04; I² statistic = 94.5%). Data from three trials contributed to the SMD for the calcipotriol versus dithranol: 0.73 (95% CI -0.55 to 2.00; I² statistic = 97.2%) (Berth Jones 1992b; Monastirli 2000; Van de Kerkhof 2006). One of these three trials found a large and statistically significant difference in favour of calcipotriol (Berth Jones 1992b); Monastirli 2000 found a significant difference in favour of dithranol, and the trial by Van de Kerkhof 2006 found no difference between the two treatments. In the light of this heterogeneity, we removed all pooling from this comparison.

Analysis 11: Vitamin D alone or in combination versus other vitamin D analogue

Our review identified three intervention-comparator contrasts for body psoriasis in this comparison: calcipotriol versus calcitriol, calcipotriol versus tacalcitol, and calcipotriol versus maxacalcitol (see Analysis 11.3 and Table 14). One between-patient trial involving 15 participants contributed PASI data for the comparison of calcipotriol and calcitriol (Bourke 1997). Treatment duration was eight weeks. Although there was a trend towards a greater effect for calcipotriol, the SMD for the PASI was not statistically significant: -1.11 (95% CI -2.22 to 0.01; I² statistic = NA).

Analysis 12: Vitamin D alone or in combination versus vitamin D + corticosteroid

Our review identified 12 intervention-comparator contrasts, involving 3 vitamin D analogues, 2 combination products, and 7 different corticosteroids (see Analysis 12.3 and Table 15). Sixteen trials involving 5703 participants contributed PASI data for 11 of these 12 intervention-comparator contrasts. Fifteen trials were between-patient, and 1 was within-patient in design (Salmhofer 2000). Treatment duration ranged from 2 to 12 weeks. We did not identify any PASI data for the comparison of calcipotriol against clobetasol propionate then calcipotriol. Overall, vitamin D plus corticosteroid appeared to be more effective than vitamin D alone: The SMD for the PASI was 0.47 (95% CI 0.34 to 0.59; I² statistic = 82.3%). However, there was considerable variation between the findings of the intervention-comparator contrasts. In light of the observed heterogeneity, we only pooled subtotals (Higgins 2011). In six of the intervention-comparator contrasts, the analysis of the PASI measure found no significant difference between the treatments. This was the case for twice-daily calcipotriol compared with the following:

 a regimen with calcipotriol applied in the morning and the night-time application of betamethasone valerate (Kragballe 1998b; Ruzicka 1998);

- clobetasone butyrate (Kragballe 1998b); or
- diflucortolone valerate (Salmhofer 2000).

Once-daily calcipotriol was no more effective than combined treatment with fluocinonide acetonide (Wozel 2001) or combined with hydrocortisone (Ortonne 2010). Lastly, no significant difference was found between twice-daily calcitriol and combined treatment with diffucortolone valerate (mornings) and calcitriol (night time) (Lee 2007).

In contrast, combined treatment with calcipotriol and betamethasone dipropionate was significantly more effective than twicedaily calcipotriol alone. This finding held for regimens involving night-time applications of betamethasone dipropionate (SMD 0.46; 95% CI 0.10 to 0.82; I² statistic = NA) and for treatment with a combined product, whether applied once daily (SMD 0.52; 95% CI 0.38 to 0.67; I² statistic = 32.3%) or twice daily (SMD 0.64; 95% CI 0.46 to 0.83; I² statistic = 73.6%). Once-daily treatment with the combined calcipotriol/betamethasone dipropionate product was also significantly more effective than once-daily calcipotriol (SMD 0.67; 95% CI 0.23 to 1.11; I² statistic = 87.4%) as well as once-daily tacalcitol (SMD 0.47; 95% CI 0.25 to 0.69; I² statistic = NA).

Analysis 13: Vitamin D alone or in combination versus other treatments: complex regimens

This comparison summarises findings on complex regimens for body psoriasis, defined here as treatment sequences that do not consist of a simple head-to-head comparison between two active treatments (see Analysis 13.3 and Table 16). PASI data were available for 11 of the 12 intervention-comparator contrasts identified. Data from 2991 participants contributed to the PASI analysis, based on findings from 8 between-patient trials. Trial duration varied between 2 and 12 weeks. Because the interventions and comparators were highly variable and two trials each contributed three pair-wise contrasts, we did not pool the data.

Six intervention-comparator contrasts for which PASI data were available found a significant difference between regimens. A sixweek course of calcipotriol was compared with two different complex regimens. Monotherapy with calcipotriol was significantly less effective than two weeks of treatment with calcipotriol and fluocinonide acetonide followed by four weeks of calcipotriol (SMD 0.66; 95% CI 0.01 to 1.32) (Wozel 2001). Calcipotriol monotherapy was also less effective than treatment with calcipotriol to which halometasone was added (SMD 1.13; 95% CI 0.64 to 1.62) (Yang 2009). Monotherapy with tacalcitol (8 weeks) was less effective than sequential treatment with a combined calcipotriol and betamethasone dipropionate product (4 weeks) followed by calcipotriol monotherapy for a further 4 weeks (SMD 0.49; 95% CI 0.31 to 0.67) (Ortonne 2004). White 2006 (P) compared three regimens, all of which included an initial phase in which participants applied combination treatment with calcipotriol and betamethasone dipropionate once a day for four weeks. The subsequent eight-week maintenance phase using placebo ointment was significantly less effective than maintenance with either twicedaily calcipotriol (SMD 0.25; 95% CI 0.10 to 0.39), or maintenance with calcipotriol on weekdays and combination treatment at weekends (SMD 0.59; 95% CI 0.45 to 0.74). When the two active maintenance regimens were compared directly, the alternating (weekday/weekend) regimen was significantly more efficacious (SMD 0.30; 95% CI 0.16 to 0.45) (White 2006 (H)).

In the remaining four intervention-comparator contrasts, we did not detect any significant difference in the PASI assessments (see Analysis 13.3). PASI assessments in the study by Kragballe 2004 found the difference between two complex regimens not to be significant (SMD 0.15; 95% CI -0.01 to 0.30). This result was on the borderline of significance and contrasted with the IAGI assessment in the same study, which was statistically significant in favour of the complex regimen (see Analysis 13.1).

Analysis 14: Vitamin D alone or in combination versus other treatment: long-term studies (> 24 weeks)

We did not identify any relevant study that provided PASI data for this analysis.

Analysis 15: Vitamin D analogues versus other treatment

This comparison incorporated all other vitamin D head-to-head comparisons of treatments for psoriasis of the body (excluding inverse psoriasis) that had not already been included (see Analysis 15.3 and Table 18). We included 12 intervention-comparator contrasts, with PASI data available for 6 of these contrasts. There were six between-patient trials and three within-patient trials, with data for 1228 participants. Trial duration ranged between 4 and 12 weeks. In light of the pharmacological diversity of the comparators, we only pooled data within subgroups (Higgins 2011). According to the PASI assessment, twice-daily calcipotriol was significantly more effective than coal tar polytherapy (SMD -0.63; 95% CI -1.06 to -0.20; I² statistic = NA) as well as propylthiouracil cream (SMD -2.24; 95% CI -3.23 to -1.25; I² statistic = NA). Twice-daily calcipotriol was not significantly more effective than coal tar (SMD -0.10; 95% CI -1.54 to 1.35; I² statistic = 92.8%), betamethasone dipropionate and salicylic acid (SMD -0.05; 95% CI -0.36 to 0.26; I² statistic = NA), or vitamin B12 cream (SMD -0.01; 95% CI -0.78 to 0.75; I² statistic = NA). The high level of heterogeneity observed in the pooled analysis of calcipotriol versus coal tar reflects contradictory findings from two studies, one finding in favour of calcipotriol (Tham 1994) and the other in favour of coal tar (Alora-Palli 2010).

When compared with once-daily dosing, twice-daily vitamin D was borderline in demonstrating a significantly better effect (SMD -0.12; 95% CI -0.25 to 0.00; I² statistic = 0%). The subgroups contributing to this result were dosing comparisons of calcipotriol (SMD -0.12; 95% CI -0.28 to 0.03; I² statistic = 0%) and combination treatment with calcipotriol and betamethasone dipropionate (SMD -0.12; 95% CI -0.32 to 0.09; I² statistic = NA).

Analysis 16: Flexural/facial psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of topical treatments for inverse or facial psoriasis (see Analysis 16.3 and Table 19). We found evidence on four treatments in this comparison: the potent steroid betamethasone valerate; the vitamin D analogue calcipotriol; and two topical calcineurin inhibitors, pimecrolimus and tacrolimus. We only identified one placebo-controlled trial evaluating tacrolimus ointment (Lebwohl 2004), but the study did not report any effectiveness data suitable for this review. However, the study did contribute data on adverse events and withdrawal rates.

PASI data were available for 3 of the 4 topical treatments for inverse psoriasis, all reported by 1 4-week between-patient study with 75 participants (Kreuter 2006 (P)). In this study, participants applied all treatments once daily. The SMD for the PASI found a statistically significant difference in favour of betamethasone valerate 0.1% (SMD -2.83; 95% CI -3.79 to -1.88) as well as calcipotriol ointment (SMD -1.08; 95% CI -1.77 to -0.40). However, PASI data on once-daily pimecrolimus cream indicated the difference relative to vehicle was not statistically significant (SMD -0.62; 95% CI -1.27 to 0.02).

Analysis 17: Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for inverse psoriasis, where vitamin D was compared with an active control (see Analysis 17.3 and Table 20). We identified five intervention-comparator contrasts. We compared four treatments with calcipotriol: once-daily betamethasone valerate, combined treatment with calcipotriol and hydrocortisone, calcitriol, and pimecrolimus. Calcitriol was compared with tacrolimus.

Two between-patient studies contributed PASI data from 464 study participants on 3 of the 5 intervention-comparator contrasts. Kreuter 2006 (H) found that calcipotriol was significantly less effective than betamethasone valerate (SMD 2.02; 95% CI 1.20 to 2.84), but not significantly different in effect to pimecrolimus (SMD -0.53; 95% CI -1.17 to 0.11). Participants in the Kreuter 2006 (H) trial applied all treatments once daily. When assessed using the PASI, Ortonne 2010 found that calcipotriol was significantly less effective on inverse psoriasis than combined treatment

with calcipotriol and hydrocortisone (SMD 0.32; 95% CI 0.12 to 0.51). This finding was consistent with the IAGI assessment in the same trial (see Analysis 17.1).

Analysis 18: Scalp psoriasis: placebo-controlled trials

No PASI data were available for this comparison.

Analysis 19: Scalp psoriasis: vitamin D alone or in combination versus other treatment

No PASI data were available for this comparison.

(d) Patient Assessment of overall Global Improvement (PAGI)/Patient Global Assessment of Disease Severity (PGA)

Analysis 1: Vitamin D analogues versus placebo

Our review included eight vitamin D analogues for body psoriasis in this comparison, with PAGI data available for three treatments (see Analysis 1.4 and Table 5). Five studies reported PAGI data, all of which were between-patient design, and 1467 participants contributed data. In all six studies, treatment duration was eight weeks. The pooled effect across the 3 studies reporting PAGI data was SMD -0.54; 95% CI -0.72 to -0.36; I² statistic = 55.5%.

Analysis 2: Corticosteroid (potent) versus placebo

Our review did not identify any study that compared potent corticosteroids against placebo and reported PAGI data.

Analysis 3: Corticosteroid (very potent) versus placebo

We included three very potent corticosteroids for body psoriasis in this comparison for this outcome (see Analysis 3.4 and Table 7). Three studies reported PAGI data for 283 participants on 2 of these 3 treatments. There was one between-patient trial (Lebwohl 2002) and two within-patient studies. In all three studies, treatment duration was two weeks. The pooled effect across both treatments for PAGI was (SMD -1.22; 95% CI -1.42 to -1.02; I² statistic = 0%). Both treatments performed statistically significantly better than placebo. PAGI data on the use of very potent steroids on the scalp are reported in Analysis 18.4.

Analysis 4: Dithranol versus placebo

Our review did not identify any study comparing dithranol against placebo and that reported PAGI data.

Analysis 5: Vitamin D combination products versus placebo

This comparison included treatment with combined calcipotriol and betamethasone dipropionate used either once or twice daily on the body (see Analysis 5.4 and Table 8). None of the studies of twice-daily treatment reported PAGI data, but one parallel-group study with 235 participants contributed data on once-daily treatment (Langley 2011 (P)). Treatment duration was eight weeks. The PAGI SMD was -0.69 (95% CI -0.98 to -0.40; I² statistic = NA). Placebo-controlled trials of combination vitamin D/steroid treatments for scalp psoriasis are reported in Analysis 18.4.

Analysis 6: Other treatment versus placebo

This comparison comprised all other treatments for body psoriasis that we did not include in the first five comparisons; therefore, we removed pooling. None of the studies assessed the same treatment, which means that findings should be interpreted with caution. In total, we included 26 treatments in this analysis (see Analysis 6.4 and Table 9). Two studies with 105 participants reported PAGI data on 2 of these 26 treatments. Both trials were between-patient studies. Treatment duration ranged from 3 to 12 weeks. According to participants' assessments (PAGI), betamethasone 17-valerate 21 acetate plus tretinoin plus salicylic acid performed significantly better than placebo (SMD -0.80; 95% CI -1.26 to -0.35), whilst kukui nut oil was no more effective than placebo (SMD 0.00; 95% CI -0.80 to 0.80). These findings were very similar to the investigators' assessments (see Analysis 6.1).

Analysis 7: Vitamin D analogues versus corticosteroid (potent)

Our review identified eight vitamin D analogue-potent corticosteroid comparisons for body psoriasis (see Analysis 7.4 and Table 10). Two studies with 738 participants reported PAGI data for 1 of these 8 comparisons. One trial was a parallel-group study (Cunliffe 1992), and one was a within-patient study (Kragballe 1991a), both with a treatment duration of six weeks. The PAGI SMD indicated that the vitamin D analogue calcipotriol was significantly more effective than betamethasone valerate (SMD -0.26; 95% CI -0.38 to -0.14; I² statistic = 0%).

Analysis 8: Vitamin D analogues versus corticosteroid (very potent)

This comparison included one vitamin D analogue and a very potent corticosteroid contrast, calcipotriol ointment versus clobetasol propionate foam for body psoriasis (see Table 11 and Analysis 8.4). One study with 42 participants reported PAGI data. Koo 2006 was a between-patient study with a treatment duration of two weeks. Findings for the participant-reported assessment (SMD 0.42; 95% CI -0.20 to 1.03) were similar, the investigators' assessment from the same study (SMD 0.19; 95% CI -0.42 to 0.80; see Analysis 8.1).

Topical treatments for chronic plaque psoriasis (Review)

Analysis 9: Vitamin D combined with corticosteroid versus corticosteroid

We found one study comparing a vitamin D analogue-steroid combination against corticosteroids for body psoriasis that reported PAGI/PGA data (see Table 12 and Analysis 9.4). This two-week between-patient study reported PAGI data for 65 participants (Koo 2006), comparing combined treatment with calcipotriol and clobetasol propionate against the very potent corticosteroid alone. According to the participants' assessment (PAGI), there was no statistically significant difference between the treatment options (SMD -0.28; 95% CI -0.80 to 0.24). This finding contrasts with the investigators' assessment (IAGI), which found in favour of combination treatment (SMD -0.69; 95% CI -1.22 to -0.15) (see Analysis 9.1).

Analysis 10: Vitamin D alone or in combination versus dithranol

This comparison considered vitamin D analogues against dithranol (see Analysis 10.4 and Table 13). We identified three intervention-comparator contrasts: calcipotriol versus dithranol, calcitriol versus dithranol, and tacalcitol versus dithranol. PAGI data were available only for the intervention-comparator contrast of calcipotriol versus dithranol. Two between-patient trials reported PAGI data for 544 participants (Berth Jones 1992b; Van de Kerkhof 2006). Treatment duration ranged from 8 to 12 weeks. The SMD for the PAGI was -0.05 (95% CI -0.90 to 0.80; I² statistic = 92.5%), indicating no statistically significant advantage for calcipotriol or dithranol according to the participants' assessment.

Analysis 11: Vitamin D alone or in combination versus other vitamin D analogue

Our review identified three intervention-comparator contrasts for body psoriasis in this comparison: calcipotriol versus calcitriol, calcipotriol versus tacalcitol, and calcipotriol versus maxacalcitol (see Analysis 11.4 and Table 14). One between-patient trial involving 250 participants contributed PAGI data for the comparison of calcipotriol and calcitriol (Ji 2008). Treatment duration was 12 weeks. The SMD for the PAGI indicated that there was no statistically significant difference between calcipotriol and calcitriol (SMD 0.04; 95% CI -0.21 to 0.29; I² statistic = NA). This study used three measures to assess the effects of calcipotriol and calcitriol. The investigators' assessment supported the participant assessment (no difference) (see Analysis 11.1), but the TSS found a significant difference in favour of calcipotriol (Analysis 11.2).

Analysis 12: Vitamin D alone or in combination versus vitamin D + corticosteroid

Our review identified 12 intervention-comparator contrasts for body psoriasis, involving 3 vitamin D analogues, 2 combination

products, and 7 different corticosteroids (see Analysis 12.4 and Table 15). Two parallel-group trials contributed PAGI data from 399 participants for 2 of these 12 intervention-comparator contrasts. Treatment duration ranged between two and eight weeks. According to the participant assessment, twice-daily calcipotriol was significantly less effective than twice-daily treatment with calcipotriol and clobetasol propionate (SMD 0.70; 95% CI 0.16 to 1.23). Treatment with tacalcitol was also less effective than treatment with a combined product containing calcipotriol and betamethasone dipropionate, when both treatments were applied once daily (SMD 0.46; 95% CI 0.24 to 0.68). The pooled SMD for these two trials (0.49; 95% CI 0.29 to 0.69; I² statistic = 0%) gave greater weight to the comparison with tacalcitol, since this was the larger trial (334 participants; Langley 2011 (H)).

Analysis 13: Vitamin D alone or in combination versus other treatments: complex regimens

This comparison summarises findings on complex regimens for body psoriasis, defined here as treatment sequences that do not consist of a simple head-to-head comparison between two active treatments (see Analysis 13.4 and Table 16). We identified 12 intervention-comparator contrasts and data from the Patient Assessment of Global Improvement (PAGI) or Patient Global Assessment of Disease Severity (PGA) were available for 7 of these. Data from 2508 participants contributed to the PAGI analysis, based on findings from 4 between-patient trials. Trial duration varied between 8 and 12 weeks. Because the interventions and comparators were highly variable and two trials each contributed three pairwise contrasts, we did not pool the data.

Four intervention-comparator contrasts for which PAGI data were available found a significant difference between regimens. Ortonne 2004 found that four weeks of treatment with a combination product of calcipotriol and betamethasone dipropionate, followed by monotherapy with calcipotriol for four weeks, was significantly more effective than monotherapy with tacalcitol (SMD 0.54; 95% CI 0.36 to 0.72). Findings from the patient-reported outcome were identical to the investigators' assessment (Analysis 13.1). The trial by White 2006 (H)/White 2006 (P) compared three regimens. All included an initial phase in which participants applied once-daily combination treatment with calcipotriol and betamethasone dipropionate for four weeks, but differed in their subsequent eight-week maintenance phase. According to the participants' assessment, maintenance therapy with twice-daily placebo ointment was significantly less effective than maintenance with either calcipotriol ointment (SMD 0.28; 95% CI 0.13 to 0.42) or maintenance with calcipotriol on weekdays and combination treatment at weekends (SMD 0.71; 95% CI 0.56 to 0.85). We compared the 2 regimens with active maintenance options directly; the alternating (weekday/weekend) approach was significantly more effective (SMD 0.44; 95% CI 0.29 to 0.58). These findings, based on the participants' assessment of severity, were aligned with those of both the investigators' assessment (Analysis 13.1) and the PASI assessment (Analysis 13.3).

A head-to-head comparison of two complex regimens found a significant difference (Kragballe 2004). Once-daily combination treatment (calcipotriol/betamethasone dipropionate) for 8 weeks followed by once-daily calcipotriol for 4 weeks was significantly less effective than a regimen consisting of combined treatment for 4 weeks, followed by 8 weeks of once-daily therapy with calcipotriol on weekdays and combined therapy at weekends (SMD 0.23; 95% CI 0.07 to 0.39).

In three of the seven intervention-comparator contrasts, no significant difference was found. Kragballe 2004 also compared 12 weeks of calcipotriol monotherapy with the 2 complex regimens described in the paragraph above. Compared with monotherapy, participants who applied once-daily combination treatment (calcipotriol/betamethasone dipropionate) for 8 weeks followed by once-daily calcipotriol for 4 weeks did not experience a significantly greater improvement in their psoriasis (SMD -0.14; 95% CI -0.30 to 0.02). Similarly, the psoriasis of participants on monotherapy with twice-daily calcipotriol did not improve significantly differently compared to those who applied once-daily combination treatment (calcipotriol/betamethasone dipropionate) for 4 weeks, followed by 8 weeks using calcipotriol once daily (weekdays) and combination therapy (weekends) (SMD 0.10; 95% CI -0.06 to 0.26).

Analysis 14: Vitamin D alone or in combination versus other treatment: long-term studies (> 24 weeks)

We did not identify any relevant study that provided PAGI data for this analysis.

Analysis 15: Vitamin D analogues versus other treatment

This comparison incorporated all other vitamin D head-to-head comparisons of treatments for psoriasis of the body (excluding inverse psoriasis) that we had not already included (see Analysis 15.4 and Table 18). We included 12 intervention-comparator contrasts, with PAGI data available for 6 of these contrasts: 3 between-patient trials and 3 within-patient trials data for 456 participants. Trial duration ranged between 4 and 12 weeks. In light of the pharmacological diversity of the comparators, we only pooled data within subgroups.

According to the participants' assessment, twice-daily calcipotriol was significantly more effective than three comparators, with evidence on each based on a single trial. Calcipotriol was more effective than coal tar both as monotherapy (SMD -1.51; 95% CI - 2.12 to -0.90; Tham 1994) and as polytherapy (SMD -0.56; 95% CI -0.99 to -0.13; Van de Kerkhof 2002a), and more effective than betamethasone dipropionate and salicylic acid (SMD -0.49; 95% CI -0.79 to -0.20; Scarpa 1994). No significant difference was evident when calcipotriol was compared to tacrolimus ointment

(Ortonne 2006), tazarotene (Tzung 2005), or vitamin B12 cream (Stuecker 2001).

Analysis 16: Flexural/facial psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of topical treatments for inverse or facial psoriasis (see Analysis 16.4 and Table 19). We found evidence on four treatments in this comparison: the potent steroid betamethasone valerate; the vitamin D analogue calcipotriol; and two topical calcineurin inhibitors, pimecrolimus and tacrolimus. We identified only one placebo-controlled trial evaluating tacrolimus ointment (Lebwohl 2004), but the study did not report any effectiveness data suitable for this review. However, the study did contribute data on adverse events and withdrawal rates.

PAGI (patient-assessment) data were available for one of the four topical treatments for inverse psoriasis. One 8-week between-patient study (Gribetz 2004) reported data from 47 participants. Relative to placebo, the SMD for the PAGI found a statistically significant difference in favour of twice-daily pimecrolimus cream (SMD -0.65; 95% CI -1.24 to -0.06).

Analysis 17: Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

We did not identify any relevant study that provided PAGI data for this analysis.

Analysis 18: Scalp psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of treatments for scalp psoriasis (see Analysis 18.4 and Table 21). We included evidence on 11 treatments in this comparison, with PAGI data available for 5 treatments. Five between-patient trials contributed data from 1875 participants. Trial duration ranged between three and eight weeks.

Four of the five treatments for scalp psoriasis that were evaluated using the Patient Global Assessment Scale were significantly more effective than placebo. The least effective treatment was calcipotriol (SMD -0.66; 95% CI -1.28 to -0.05; I² statistic = 74.5%), and the most effective was betamethasone dipropionate (SMD -1.23; 95% CI -1.43 to -1.03; I² statistic = NA). Effects for the very potent steroid amcinonide (SMD -0.97; 95% CI -1.33 to -0.61 I² statistic = NA) and combination treatment with calcipotriol and betamethasone dipropionate (SMD -1.00; 95% CI -1.79 to -0.22; I² statistic = 93.2%) fell between these two extremes. There were no data suitable for pooling across subcategories.

Only for ciclopirox olamine shampoo was the difference relative to placebo found to be non-significant: SMD -0.11 (95% CI - 0.86 to 0.64; I² statistic = NA).

Analysis 19: Scalp psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for scalp psoriasis in which one of the interventions was a vitamin D product (used either as monotherapy or in combination with another product) (see Analysis 19.4 and Table 22).

We identified six intervention-comparator contrasts, and PAGI data were available for four of these contrasts (there were no participant-assessed outcomes data for the comparison of calcipotriol against either clobetasol propionate or against coal tar polytherapy). All studies were parallel-group in design (between-patient). Six studies contributed PAGI data from 3742 participants, and trial duration ranged from 4 to 8 weeks.

Based on the participant assessment scores, calcipotriol was significantly less effective than betamethasone dipropionate (SMD 0.56; 95% CI 0.31 to 0.81; I² statistic = 82.8%), betamethasone valerate (SMD 0.41; 95% CI 0.22 to 0.59; I² statistic = NA), and combination treatment with calcipotriol and betamethasone dipropionate (SMD 0.84; 95% CI 0.61 to 1.08; I² statistic = 81.5%). Combination treatment (calcipotriol/betamethasone dipropionate) was significantly more effective than betamethasone dipropionate alone (SMD -0.17; 95% CI -0.25 to -0.09; I² statistic = 0%).

(e) Combined end point (IAGI/TSS/PASI/PAGI)

Analysis 1: Vitamin D analogues versus placebo

Our review included eight vitamin D analogues for body psoriasis in this comparison (see Analysis 1.5 and Table 5). Thirty studies, involving 4986 participants, contributed data. Eighteen trials were between-patient design, and 12 were within-patient studies. Treatment duration ranged from 3 to 12 weeks. Six studies had adequately concealed treatment allocation. The pooled effect across all 30 studies was -0.90 (95% CI -1.07 to -0.72; I2 statistic = 87.5%), which equates to 1.03 on a 6-point IAGI scale. However, the summary statistic conceals considerable variation between treatments. In two treatments, the effect was not statistically significantly different to placebo (twice-daily calcipotriol plus occlusion for 2 weeks followed by 4 weeks with no treatment, and oncedaily becocalcidiol). In treatments that were significantly more effective than placebo, the effect ranged from -0.67 (twice-daily becocalcidiol) to -1.66 (once-daily paricalcitol); on a 6-point IAGI scale, these effects translate into 0.80 and 1.91 points, respectively. However, given the high level of heterogeneity found in the metaanalysis, it may be more informative to look at individual products within the class, rather than treating this class as a single group. We removed the pooling across the class, so that we only pooled subtotals.

We reported placebo-controlled trials of vitamin D products for scalp psoriasis (Analysis 18.5) and inverse psoriasis (Analysis 16.5) elsewhere.

Sensitivity analyses

We explored differences in within-patient and between-patient designs using one-way sensitivity analysis (Table 5). The SMD for the 12 within-patient studies (600 participants) was -1.11 (95% CI -1.58 to -0.64; 1² statistic = 91.5%). This translates into a change of 1.27 on a 6-point IAGI scale, slightly larger than the effect size for all studies. The SMD for the 18 between-patient studies (4386 participants) was -0.80 (95% CI -0.96 to -0.63; 1² statistic = 83.2%). This translates into a change of 0.91 on a 6-point IAGI scale, slightly smaller than the effect size for the within-patient studies.

We also used sensitivity analysis to explore the impact of varying the correlation coefficient (rho) for the 12 within-patient studies. As no trial in the review reported this statistic, we tested values of 0, 0.25, 0.50, and 0.75. Varying the value of rho for the within-patient studies had no significant effect on the findings: The pooled SMD ranged from -0.85 (95% CI -1.00 to -0.71; I² statistic = 87.8%; 30 studies), when we assumed the correlation to be zero, to -0.91 (95% CI -1.07 to -0.75; I² statistic = 93.2%; 30 studies), when we assumed the correlation to be 0.75.

The study by Perez 1996 comparing calcitriol and placebo reported large and statistically significant differences for both TSS and IAGI outcomes. In both outcomes, the magnitude of the effect was the largest across all comparisons and treatments. Perez 1996 was a 10-week within-patient study involving 84 participants. The study included people with severe disease (mean TSS at baseline: 7.6 on a 10-point scale, with at least 10% of body surface area affected) who had previously had an unsatisfactory response to at least 1 previous treatment including topical steroids, UVB, PUVA, and methotrexate. The dramatic improvement observed in the intervention group is difficult to interpret in the context of findings from other trials, and we explored the impact of removing this study in a sensitivity analysis. When we removed Perez 1996 from the pooled analysis for calcitriol, the effect size was smaller but still statistically significantly different from placebo: SMD with Perez 1996 was -0.92 (95% CI -1.54 to -0.29; I² statistic = 94.9%; 7 studies), and SMD without Perez 1996 was -0.60 (95% CI -0.78 to -0.41; I² statistic = 30%; 6 studies). On a 6-point IAGI scale, these effect sizes equate to 1.05 and 0.68, respectively (Table 5). The trials of calcipotriol varied by dose, treatment duration, and dosing frequency; where trials reported more than one dose (Kragballe 1988b) or type of vehicle (Harrington 1996a), we estimated the weighted mean and standard deviation across the trial. The effect size for twice-daily regimens was -1.02 (95% CI -1.23 to -0.82 I² statistic = 73.5%; 13 studies); this equates to 1.18 on a 6-point IAGI scale. For once-daily calcipotriol, the corresponding figures were -0.76 (95% CI -1.13 to -0.40; I2 statistic = 82.3%; 4 studies) and 0.87 on a 6-point IAGI scale. Therefore, both dosing frequencies were more effective than placebo, but once-daily applications had a smaller effect (Table 5).

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Analysis 2: Corticosteroid (potent) versus placebo

Our review identified 10 potent corticosteroids for body psoriasis in this comparison (see Analysis 2.5 and Table 6). The treatments included three betamethasone dipropionate regimens, but we identified no effectiveness evidence for budesonide and no study reporting PAGI data. Therefore, 13 studies involving 2216 participants contributed data on 9 of the 10 treatments. Eleven trials were between-patient design, and two were within-patient studies. Treatment allocation was adequately concealed in four studies. Treatment duration ranged from 2 to 12 weeks. Participant numbers for individual studies ranged from 9 (Wortzel 1975 (2)) to 633 (Kaufmann 2002 (P)). The pooled effect across all 13 studies was -0.89 (95% CI -1.06 to -0.72; I² statistic = 65.1%). This equates to 1.02 on a 6-point IAGI scale. With the exception of diflorasone diacetate (Lane 1983), all potent corticosteroids were significantly more effective than placebo at the 5% level of significance. We reported elsewhere placebo-controlled trials of potent corticosteroid products for scalp psoriasis (Analysis 18.5) and inverse psoriasis (Analysis 16.5).

Sensitivity analyses

We explored differences in within-patient and between-patient designs using one-way sensitivity analysis (Table 6). The SMD for the 2 within-patient studies was -1.33 (95% CI -1.78 to -0.89; I^2 statistic = 0%). This translates into a change of 1.53 on a 6point IAGI scale, which is larger than the effect for all studies. However, as just 48 participants contributed data to this analysis, robust inferences cannot be drawn. The SMD for the 11 betweenpatient studies (2168 participants) was -0.85 (95% CI -1.03 to -0.67; I² statistic = 66.7%). This translates into a change of 0.98 on a 6-point IAGI scale, slightly smaller than the effect for all studies. We also used sensitivity analysis to explore the impact of varying the correlation coefficient (rho) for the 2 within-patient studies. As no trial in the review reported this statistic, we tested values of 0, 0.25, 0.50, and 0.75. Varying the value of rho for the withinpatient studies had no significant effect on the findings: the pooled SMD ranged from -0.89 (95% CI -1.06 to -0.72; I² statistic = 77.7%; 14 studies), when we assumed the correlation was to be zero, to -0.91 (95% CI -1.08 to -0.74; I2 statistic = 80.2%; 13 studies), when we assumed the correlation to be 0.75.

Analysis 3: Corticosteroid (very potent) versus placebo

Our review identified three very potent corticosteroids for body psoriasis in this comparison (see Analysis 3.5 and Table 7), but effectiveness data were not available for one of these treatments (halcinonide). Ten studies reported data for 1264 participants. There were seven between-patient trials and three within-patient studies. Treatment duration ranged from two to four weeks. In no study was treatment allocation adequately concealed. The SMD across the 2 treatments was -1.56 (95% CI -1.87 to -1.26; I² statistic = 81.7%; 10 studies). This equates to 1.80 points on a 6-point IAGI scale. Both clobetasol propionate (SMD -1.65; 95% CI -2.10 to -1.20; I² statistic = 86.3%; 7 studies) and halobetasol (SMD -1.36; 95% CI -1.65 to -1.07 I² statistic = 47.1%; 3 studies) performed significantly better than placebo.

We reported elsewhere placebo-controlled trials of very potent corticosteroid products for scalp psoriasis (Analysis 18.5).

Sensitivity analyses

We explored differences in within-patient and between-patient designs using one-way sensitivity analysis (Table 7). The SMD for the 3 within-patient studies (229 participants) was -1.52 (95% CI -2.02 to -1.02; I² statistic = 79.3%). This translates into a change of 1.74 on a 6-point IAGI scale. The SMD for the 7 between-patient studies (1035 participants) was -1.58 (95% CI -1.99 to -1.17; I² statistic = 84.4%). This translates into a change of 1.81 on a 6-point IAGI scale.

We also used sensitivity analysis to explore the impact of varying the correlation coefficient (rho) for the 3 within-patient studies. As no trial in the review reported this statistic, we tested values of 0, 0.25, 0.50, and 0.75. Varying the value of rho for the within-patient studies had no significant effect on the findings: The pooled SMD ranged from -1.52 (95% CI -1.80 to -1.24; I² statistic = 81.6%; 10 studies), when we assumed the correlation to be zero, to -1.55 (95% CI -1.80 to -1.29; I² statistic = 85.9%; 10 studies), when we assumed the correlation to be 0.75.

Analysis 4: Dithranol versus placebo

This comparison considered dithranol against placebo (see Analysis 4.5 and Table 23). Three within-patient trials reported data for 47 participants. The only outcome reported by these trials was the TSS, so results for the combined end point are identical to the TSS results section. The types of treatment comprised the following: dithranol 0.1% in a carbamide (17% urea) base twice daily (Buckley 1978), dithranol in aqueous gel (dose titration 0.1% to 2.0%) twice daily (Grattan 1997 (P)), and dithranol 2% ointment 'one minute therapy' once daily (Jekler 1992).

Treatment duration ranged from three to eight weeks. In all three studies, the adequacy of the concealment of treatment allocation was unclear. The SMD for the combined end point was -1.06 (95% CI -1.66 to -0.46; I² statistic = 37.4%; 3 studies), which equates to 1.22 on a 6-point IAGI scale. All three trials found a statistically significant effect in favour of dithranol.

Sensitivity analysis

We used sensitivity analysis to explore the impact of varying the correlation coefficient (rho) for the 3 within-patient studies. As no

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trial in the review reported this statistic, we tested values of 0, 0.25, 0.50, and 0.75. Varying the value of rho for the within-patient studies had no significant effect on the findings: The pooled SMD ranged from -0.98 (95% CI -1.56 to -0.41; I² statistic = 13.9%; 3 studies), when we assumed the correlation to be zero, to -1.17 (95% CI -1.81 to -0.52; I² statistic = 78.5%), when we assumed the correlation to be 0.75.

Analysis 5: Vitamin D combination products versus placebo

This comparison included treatment with combined calcipotriol and betamethasone dipropionate used either once or twice daily on the body (see Analysis 5.5 and Table 8). As all five studies reported IAGI data, results for the combined end point are identical to those in Analysis 5.1. Five parallel-group studies with 2058 participants contributed data. Treatment duration ranged from four to eight weeks. Three trials adequately concealed treatment allocation. The SMD for the combined end point across treatments was -1.44 (95% CI -1.76 to -1.12; I² statistic = 89.4%; 5 studies), with twice-daily combination treatment (SMD -1.90; 95% CI -2.09 to -1.71; 2 studies) achieving a significantly larger effect than oncedaily treatment (SMD -1.21; 95% CI -1.50 to -0.91; 3 studies). On a 6-point IAGI scale, these equate to improvements of 2.18 and 1.39 points, respectively.

However, the PASI assessments differed from the IAGI: although twice-daily combination treatment achieved a larger effect than once-daily treatment (SMD -1.41 and -1.14, respectively), the difference was not statistically significant (see Analysis 5.3).

In Analysis 18.5, we reported placebo-controlled trials of combination vitamin D/steroid treatments for scalp psoriasis.

Analysis 6: Other treatment versus placebo

This comparison comprised all other treatments for psoriasis of the body that we did not include in the first five comparisons; therefore, we removed pooling. None of the studies assessed the same treatment, which means that findings should be interpreted with caution.

In total, we included 26 treatments in this analysis (see Analysis 6.5 and Table 9). Twenty-six studies, with 1450 participants, reported data on 25 of these 26 treatments. The study of omega-3-polyunsaturated fatty acids ointment (Henneicke-v. Z. 1993) did not report usable effectiveness data, but did contribute data to the analysis of withdrawals. Twelve trials were between-patient design, and 14 were within-patient studies. Treatment duration ranged from 2 to 12 weeks. Two studies adequately concealed treatment allocation (Levine 2010 (P); Stutz 1996).

As assessed by the combined end point, just over half the treatments (13/25) performed statistically significantly better than placebo: aloe vera cream, anti-IL-8 monoclonal antibody cream, betamethasone 17-valerate 21-acetate plus tretinoin plus salicylic acid, combined treatment with calcipotriene and nicotinamide, fish oil plus occlusion, herbal skin care products, indigo naturalise 1.4% ointment, *Mahonia aquifolium*, methotrexate gel, mycophenolic acid ointment, PTH (1-34) in Novasome cream®, tazarotene, and theophylline 1% ointment. The effect size (SMD) for the combined end point ranged from -0.48 (Levine 2010 (P); calcipotriene combined with nicotinamide) to -2.96 (Maier 2004; herbal skin care products), or 0.55 to 3.40, respectively, on a 6point IAGI scale.

The trial of herbal skin care products enrolled just 34 participants, of which 24 contributed data (Maier 2004), so this study is classed as having a high risk of attrition bias (Figure 3; see Risk of bias in included studies). The results of this study were published as an abstract, and our searches failed to locate a full publication. Therefore, findings should be treated with caution.

In the remaining 12 treatments, the difference relative to placebo using the combined end point was not statistically significant. These included topical caffeine, Dead Sea salts emollient lotion, hexafluoro-1,25-dihydroxyvitamin D3, kukui nut oil, NGmonomethyl-L-arginine (L-NMMA) cream, nicotinamide 1.4%, oleum horwathiensis, platelet aggregation activating factor (PAF), polymyxin B cream, topical sirolimus, topical tacrolimus, and tar. In two treatments, findings by different outcomes were inconsistent. When assessed by the TSS, hexafluoro-1,25-dihydroxyvitamin D3 was significantly more effective than placebo (-1.13; 95% CI -1.91 to -0.35) (Analysis 6.2), whereas the difference assessed by the IAGI was non-significant (-0.62; 95% CI -1.35 to 0.12) (Analysis 6.1). We based these findings on a single trial (Durakovic 2001). Similarly, the difference for oleum horwathiensis was statistically significant using the TSS (-0.77; 95% CI -1.40 to -0.14) (Analysis 6.2) whereas the difference assessed by the IAGI was not (-0.02; 95% CI -0.63 to 0.58) (Analysis 6.1). We also based findings for this treatment on a single trial (Lassus 1991).

We reported elsewhere placebo-controlled trials of other treatments for scalp psoriasis (Analysis 18.5) and inverse psoriasis (Analysis 16.5).

Analysis 7: Vitamin D analogues versus corticosteroid (potent)

Our review identified eight vitamin D analogue-potent corticosteroid comparisons for body psoriasis for this comparison (see Analysis 7.5 and Table 10). Fourteen studies with 3542 participants reported data for these 8 intervention-comparator contrasts. Nine trials were between-patient design, and five were within-patient studies. Treatment duration ranged from three to eight weeks. One study adequately concealed treatment allocation (Papp 2003 (H)).

The combined end point SMD indicated that, overall, there was no statistically significant difference between vitamin D and potent corticosteroids: SMD 0.11 (95% CI -0.07 to 0.30; I² statistic = 85.6%; 14 studies). There was however substantial heterogeneity and variation in effect underlying this summary statistic. Therefore, we only pooled subtotals.

The difference between the vitamin D analogue and the potent corticosteroid was not statistically significant for four interventioncomparator contrasts (calcipotriol versus betamethasone valerate, calcipotriol versus desoxymetasone, calcitriol versus betamethasone dipropionate, calcitriol versus betamethasone valerate). In the comparison of calcipotriol and betamethasone valerate, the nonsignificant result was borderline (SMD -0.12; 95% CI -0.26 to 0.02; I² statistic = 41.6%; 4 studies). When assessed using the TSS (one study), PASI (four studies), and PAGI (two studies), calcipotriol was significantly more effective than betamethasone valerate (see Analysis 7.2, Analysis 7.3, and Analysis 7.4). There were similar inconsistencies between the outcome assessments for the comparison of calcitriol with betamethasone dipropionate, all of which we based on a single study (Camarasa 2003). According to the TSS and PASI, calcitriol was significantly less effective than betamethasone dipropionate, but the IAGI indicated that the difference was not significant (see Analysis 7.1).

In one intervention-comparator contrast (calcipotriol versus fluocinonide), the vitamin D analogue was significantly more effective than potent corticosteroid: SMD -0.58 (95% CI: -0.99 to -0.18; I² statistic = NA). This effect is equivalent to a gain of approximately two-thirds of a point (0.64) on a 6-point IAGI scale. In three intervention-comparator contrasts, the potent corticosteroid was significantly more effective than the vitamin D analogue (calcipotriol versus betamethasone dipropionate, calcipotriol versus diflorasone diacetate, tacalcitol versus betamethasone valerate). The SMD effect sizes for these were 0.43, 0.27, and 0.41, respectively, equivalent to improvements of 0.47, 0.30, and 0.45 on a 6-point IAGI scale.

We reported elsewhere trials comparing vitamin D and potent corticosteroids for scalp psoriasis (Analysis 19.5) and inverse psoriasis (Analysis 17.5).

Sensitivity analyses

We explored differences in within-patient and between-patient designs using one-way sensitivity analysis (Table 10). In both types of design, the difference between vitamin D and potent corticosteroid was not statistically significant, but we associated the pooled results with substantial levels of heterogeneity (Higgins 2011). The SMD for the 5 within-patient studies (554 participants) was 0.17 (95% CI -0.20 to 0.54; I² statistic = 81.9%). The SMD for the 9 between-patient studies (2988 participants) was 0.10 (95% CI -0.11 to 0.31; I² statistic = 84.9%).

We also used sensitivity analysis to explore the impact of varying the correlation coefficient (rho) for the 5 within-patient studies. As no trial in the review reported this statistic, we tested values of 0, 0.25, 0.50, and 0.75. Varying the value of rho for the within-patient studies had no significant effect on the findings: The pooled SMD ranged from 0.10 (95% CI -0.08 to 0.28; I² statistic = 90.5%; 14 studies), when we assumed the correlation to be zero, to 0.12 (95% CI -0.06 to 0.30; I^2 statistic = 94.3%), when we assumed the correlation to be 0.75.

Analysis 8: Vitamin D analogues versus corticosteroid (very potent)

This comparison considered vitamin D analogues against very potent corticosteroids for body psoriasis (see Analysis 8.5 and Table 11). We found data on one intervention-comparator contrast: calcipotriol versus clobetasol propionate. Two between-patient trials reported data for 82 participants. Treatment duration ranged between two and six weeks. The adequacy of treatment allocation was unclear in both studies. The SMD for the combined end point indicated that there was no statistically significant difference between the vitamin D analogue and the very potent corticosteroid: -0.06 (95% CI -0.57 to 0.44; I² statistic = 25.7%; 2 studies). We reported elsewhere trials comparing vitamin D and very potent corticosteroids for scalp psoriasis (Analysis 19.5).

Analysis 9: Vitamin D combined with corticosteroid versus corticosteroid

This comparison considered combined treatment with vitamin D analogues and corticosteroids against potent or very potent corticosteroid for body psoriasis (see Analysis 9.5 and Table 12). Five parallel-group studies, ranging in treatment duration from 2 to 8 weeks, provided data on 2113 participants. The adequacy of the concealment of treatment allocation was unclear in all five trials. Effectiveness data were available for three contrasts:

• calcipotriol plus betamethasone dipropionate versus betamethasone dipropionate;

• calcipotriol plus betamethasone dipropionate versus clobetasol propionate; and

• calcipotriol plus clobetasol propionate versus clobetasol propionate.

Overall, combination treatment was more effective than monotherapy with corticosteroids: SMD -0.26 (95% CI -0.52 to -0.00; I² statistic = 84.4%; 5 studies). This effect is equivalent to a change of 0.29 on a 6-point IAGI scale. However, the pooled finding masks important diversity between the three interventioncomparator contrasts. When we compared combination treatment with calcipotriol and betamethasone dipropionate with a potent steroid (betamethasone dipropionate alone), combination treatment was more effective (SMD -0.40; 95% CI -0.52 to -0.27; I² statistic = 41.8%; 3 studies). However, the same combination treatment was significantly less effective than monotherapy with a very potent steroid (clobetasol propionate) (SMD 0.45; 95% CI 0.09 to 0.81; I² statistic = NA). Calcipotriol in combination with clobetasol propionate however was more effective than the very potent steroid used alone (SMD -0.69; 95% CI -1.22 to -0.15; I² statistic = NA). Therefore, we removed pooling for this analysis,

but the findings demonstrate that combination treatment with vitamin D analogue and a corticosteroid is more effective than the same steroid used as a monotherapy.

We reported elsewhere trials comparing combination therapy (with vitamin D and potent corticosteroids) against potent steroids for scalp psoriasis (Analysis 19.5).

Analysis 10: Vitamin D alone or in combination versus dithranol

This comparison considered vitamin D analogues against dithranol for body psoriasis (see Analysis 10.5 and Table 13). We identified three intervention-comparator contrasts: calcipotriol versus dithranol, calcitriol versus dithranol, and tacalcitol versus dithranol. Seven between-patient trials and one within-patient trial (Grattan 1997 (H)) reported data for 1284 participants. We considered concealment of treatment allocation to be adequate in one trial (Van de Kerkhof 2006). Treatment duration ranged from 4 weeks to 12 weeks. In addition, there was some variation in the dithranol regimens employed by trials and in the baseline severity of trial participants. These factors may explain the high level of heterogeneity found in the pooled results.

The SMD for the combined end point was 0.09 (95% CI -0.44 to 0.63; I² statistic = 94.9%; 8 studies), indicating that there was no evidence of a statistically significant difference in effect between vitamin D analogues and dithranol. The pooled findings for calcipotriol against dithranol (SMD 0.07; 95% CI -0.57 to 0.71; I² statistic = 95.7%; 6 studies) and tacalcitol versus dithranol (SMD -0.18; 95% CI -0.60 to 0.25; I² statistic = NA) were aligned with, and drove, this result. However, the high levels of heterogeneity mean that findings merit closer scrutiny. Six studies contributed data from 1086 participants to the comparison of calcipotriol and dithranol. In three of these studies, calcipotriol performed significantly better than dithranol; two trials found significant differences in favour of dithranol; and one study found no significant difference between the treatments (see Analysis 10.5). A single study of 114 participants compared calcitriol against dithranol (Hutchinson 2000) and found a statistically significant difference in favour of dithranol (SMD 0.51; 95% CI 0.14 to 0.88; I² statistic = NA). This equates to just over half a point (0.56 of a)point) improvement on a 6-point IAGI scale. However, the same study found no significant difference between the treatments in the assessment using the TSS (Analysis 10.2) or the PASI (Analysis 10.3).

In light of the considerable level of observed variation within and between studies (Higgins 2011), we removed pooling from this comparison.

Analysis 11: Vitamin D alone or in combination versus other vitamin D analogue

Our review identified three intervention-comparator contrasts for body psoriasis in this comparison: calcipotriol versus calcitriol, calcipotriol versus tacalcitol, and calcipotriol versus maxacalcitol (see Analysis 11.5 and Table 14). Four trials involving 513 participants contributed data. One trial was within-patient (Barker 1999 (H)); three were between-patient in design; and no trials demonstrated the concealment of treatment allocation to be adequate. Treatment duration ranged from 8 to 12 weeks.

The SMD for the combined end point indicated that there was no statistically significant difference between the treatments: -0.17 (95% CI -0.62 to 0.27; I² statistic = 78.5%; 4 studies). When we considered individual intervention-comparator contrasts using the combined end point, calcipotriol was more effective than tacalcitol (SMD -0.47; 95% CI -0.73 to -0.21; I² statistic = NA), which equates to an improvement of 0.52 on a 6-point IAGI scale. However, there was no statistically significant difference between calcipotriol and calcitriol (SMD -0.41; 95% CI -1.46 to 0.64; I² statistic = 72.3%; 2 studies) or between calcipotriol and maxacalcitol (SMD 0.43; 95% CI -0.12 to 0.98; I² statistic = NA). Participants in all trials applied treatments twice daily, with the exception of tacalcitol, which was applied once daily (Veien 1997).

Sensitivity analysis

The trial by Ji 2008 used three outcome measures to assess the effects of calcipotriol and calcitriol. Both the investigators' assessment (Analysis 11.1) and the participants' assessment found no difference between the two vitamin D products (Analysis 11.4), but the TSS found a statistically significant difference in favour of calcipotriol (Analysis 11.2). As findings for Ji 2008 varied depending on the outcome measure used, we tested the impact of using TSS data for the combined end point instead of IAGI data. The sensitivity analysis demonstrated that findings were robust, both for the SMD for the combined end point for all treatments (SMD -0.28; 95% CI -0.66 to 0.10; I² statistic = 70.6%; 4 studies) and for the comparison of calcipotriol and calcitriol (SMD -0.52; 95% CI -1.19 to 0.15; I² statistic = 44.9%; 2 studies).

Analysis 12: Vitamin D alone or in combination versus vitamin D + corticosteroid

Our review identified 12 intervention-comparator contrasts for body psoriasis, involving 3 vitamin D analogues, 2 combination products, and 7 different corticosteroids (see Analysis 12.5 and Table 15). Seventeen trials involving 5856 participants contributed data. Sixteen trials were between-patient, and 1 was within-patient in design (Salmhofer 2000). Treatment duration ranged from 2 to 12 weeks. Four trials adequately concealed treatment allocation, but concealment was inadequate in one trial (in the other trials, insufficient data were reported to allow an assessment of adequacy).

Overall, vitamin D plus corticosteroid was more effective than vitamin D alone: The SMD for the combined end point was 0.46 (95% CI 0.33 to 0.59; I^2 statistic = 83.3%; 17 studies), which

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translates into half of 1 point (0.50) on a 6-point IAGI scale. The substantial level of heterogeneity reflects not only different dosing schedules of the intervention and comparator products, but also the different potency of the corticosteroids and the different vitamin D analogues evaluated; we therefore removed pooling (Higgins 2011).

Twice-daily calcipotriol was significantly less effective than combination treatment with betamethasone dipropionate. This finding held whether participants applied the corticosteroid separately at night time (SMD 0.56; equivalent to 0.61 on a 6-point IAGI scale) or as a combined product used once daily or twice daily (SMD 0.43 and 0.66, respectively, translating into improvements of 0.48 and 0.73 points on a 6-point IAGI scale). Twice-daily calcipotriol was also significantly less effective than combination treatment with clobetasone butyrate (SMD 0.27; 95% CI 0.05 to 0.48; I² statistic = NA; 0.29 of a point on a 6-point IAGI) or clobetasol propionate (SMD 0.88; 95% CI 0.34 to 1.42; I² statistic = NA; 0.97 of a point on a 6-point IAGI scale). Once-daily combination treatment with calcipotriol and betamethasone dipropionate was significantly more effective than once-daily treatment with either calcipotriol (SMD 0.66; 95% CI 0.31 to 1.02; I² statistic = 80.9%; 2 studies) or tacalcitol (SMD 0.48; 95% CI 0.26 to 0.70; I^2 statistic = NA).

In none of the 12 intervention-comparator contrasts was the vitamin D analogue significantly more effective than combined vitamin D plus corticosteroid. However, in five instances, there was no significant difference between vitamin D and the comparator (combination treatment). These included twice-daily calcipotriol against combination treatment with betamethasone valerate or diflucortolone valerate, once-daily calcipotriol against combination treatment with fluocinonide acetonide or hydrocortisone, and twice-daily calcitriol against combination treatment with diflucortolone valerate.

We reported elsewhere trials comparing vitamin D and combination therapy with vitamin D/potent corticosteroids for scalp psoriasis (Analysis 19.5).

Analysis 13: Vitamin D alone or in combination versus other treatments: complex regimens

This comparison summarises complex regimens for body psoriasis, defined here as treatment sequences that do not consist of a simple head-to-head comparison between two active treatments (see Analysis 13.5 and Table 16). Using the combined end point, data were available for all 12 intervention-comparator contrasts (Figure 4). Data were available from 2936 participants from 8 between-patient trials and 1 within-patient trial (Austad 1998). Trial duration varied between 2 and 12 weeks. One trial adequately concealed treatment allocation. As the interventions and comparators were highly variable and because two trials each contributed three pair-wise contrasts, we did not pool the data.

Figure 4. Forest plot of comparison: 13 Vitamin D alone or in combination vs. other treatments: complex regimens, outcome: 13.5 Combined end point (IAGI/TSS/PASI/PAGI).

Std. Mean Difference tudy or Subgroup V, Random, 95% Cl	Std. Mean Difference IV, Random, 95% Cl
5.5.1 Calcipotiol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) applie 2004 -012 {0.23, 0.04} -012 {0.23, 0.04} berogenety: Not applicable Entorwealt #etc. 2 = 14 (8 P = 014) Entorwealt #etc. 2 = 14 (8 P = 014)	-
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5.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (wk)/ & combined calcipotriol + BMD (wks) dpabile 2004 0.13 [0.04, 0.29] drong emety, Nt applicable terrogenety, Nt applicable (15 or overail (BMC 2 = 1.5 (P = 0.15)	•
5.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks) etaal 1980 0.0 (0 11, 1 02) torogeneek, Not applicable torogeneek, Not applicable torovenai (efter 2 = 2 00 (7 = 0.05)	*
5.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks) zei 2001 0.6 6 (0.01, 1.32] Brorgenetik, Not applicable troroveni (effect 2 = 1.8 gP = 0.05)	*
5.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (wky), halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks) ng 200 0 4.1 (4.0.5, 0.8) biotral (95% C) 4.1 (4.0.5, 0.6) terrogeneek rol applicable s for overal (#dec 2.2 - 176 (P = 0.0))	*
5.57 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12) the fractional (5 to 4 wks); then calcitriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12) the fractional (5 to 4 wks); then calcitriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12) the fractional (5 to 4 wks); then calcitriol twice daily (to wk 12) vs. calcitriol (ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12) the fractional (5 to 4 wks); then calcitriol twice daily (to wk 12) the fractional (5 to 4 wks); then calcitriol (5 to	*
S. Combined calciplorfiel - BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calciporfiel + BMD (4 wks); then calciporfiel ointment twice daily (8 wks) the 2006 (P) 0.27 (0.12, 0.41) theragenetic, Not applicable terropenetic, 2 36 (F = 0.003)	₹
5.9 Combined calcipotriol + BMD (4 wiss); then placebo ointment twice daily (8 wiss) vs. combined calcipotriol + BMD (4 wiss); then calcipotriol (wiby)+ combined calcipotriol + BMD (wio) (8 wiss) the 2006 (P) 0.51 (0.37, 0.66] terrogeneity. Not applicable terrogeneity. Cost plass 6.55 (P < 20001)	₹
5.10 Combined calcipotriol + BMD (4 wks); then calcipotriol eintment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks) the 2006 (+) 0.25 (0.11, 0.0] total (45%). (C) 0.25 (0.11, 0.0] total (45\%). (C	
5.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (widy) & combined calcipotriol + BMD (w/e) (8 wks) gballe 2004 0.24 (0.00, 0.40) terogenety. Not applicable terogenety. Not applicable	•
5.12 Tracalctol (8 w/s) vs. combined catcipotriol • BMD (4 w/s); then catcipotriol (4 w/s) torne 2004 0.54 (0.36, 0.72] torogenetic Not applicable tor oreal effect = 2.5 & 66 (* 0.00001)	
set for subkrown diffurence: "NF-55-51 vf-11 // v // 0.00011 VF-0.0 %	-2 -1 0 1 2 Favours vitamin D regimen Favours complex regimen

Test for subaroup differences: Chi^p = 55.51, df = 11 (P < 0.00001), P = 80.2%

Seven intervention-comparator contrasts found a significant difference between regimens. Six weeks' monotherapy with twicedaily calcipotriol was less effective than clobetasol propionate (2 weeks) followed by 4 weeks' monotherapy with calcipotriol (SMD 0.60; 95% CI 0.18 to 1.02) (Austad 1998). A similar effect was evident when calcipotriol monotherapy (once daily for two weeks, then twice daily for four weeks) was compared with two weeks of treatment with calcipotriol (mornings) and fluocinonide acetonide (evenings), followed by 4 weeks' monotherapy with calcipotriol (SMD 0.66; 95% CI 0.01 to 1.32) (Wozel 2001). These benefits translated into improvements of 0.66 and 0.72 (i.e. around twothirds of a point) on a 6-point IAGI scale.

The 12-week trial by White 2006 (H)/White 2006 (P) compared 3 regimens; all included an initial phase in which participants applied once-daily combination treatment with calcipotriol and betamethasone dipropionate for 4 weeks, but differed in their subsequent 8-week maintenance phase. Maintenance therapy with twice-daily placebo ointment was significantly less effective than maintenance with either calcipotriol ointment (SMD 0.27; 95% CI 0.12 to 0.41) or maintenance with calcipotriol on weekdays and combination treatment at weekends (SMD 0.51; 95% CI 0.37 to 0.66). When we directly compared the two regimens with active maintenance options, the alternating (weekday/weekend) approach was significantly more effective (SMD 0.26; 95% CI 0.11 to 0.40). The benefits of these 3 regimens equate to an improvement of 0.29, 0.56, and 0.28 of a point, respectively, on a 6-point IAGI scale. These findings, based on the investigators' assessment of severity, were aligned with those of both the PASI assessment (Analysis 13.3) and the participants' assessment (PAGI) (Analysis 13.4).

Ortonne 2004 demonstrated that four weeks of treatment with a combination product of calcipotriol and betamethasone dipropionate followed by monotherapy with calcipotriol for four weeks was significantly more effective than monotherapy with tacalcitol (SMD 0.54; 95% CI 0.36 to 0.72) (0.59 on a 6-point IAGI scale).

Findings from the investigators' assessment (Analysis 13.1) were aligned with the PASI assessment (Analysis 13.3) and the participants' assessment (Analysis 13.4).

Kragballe 2004 compared 12 weeks of calcipotriol monotherapy with 2 complex regimens, both of which involved different sequences of combination therapy (calcipotriol/betamethasone dipropionate) and calcipotriol. When we directly compared these two complex regimens, 8 weeks of combination therapy followed by 4 weeks of once-daily calcipotriol was significantly less effective than a routine involving 4 weeks of combination therapy followed by 8 weeks of once-daily calcipotriol on weekdays and combination therapy at the weekend (SMD 0.24; 95% CI 0.08 to 0.40, equivalent to 0.27 (¼ of 1 point) on a 6-point IAGI scale). This finding was based on the investigators' assessment (IAGI), which was very similar to the participants' (PAGI) assessment (Analysis 13.4). However, the PASI found a non-significant trend in favour of the weekday/weekend regimen (Analysis 13.3).

In five intervention-comparator contrasts, we found no significant difference. A single study reported two of the non-significant intervention-comparator contrasts: Kragballe 2004 compared 12 weeks of calcipotriol monotherapy with each of the 2 complex regimens described in the paragraph above. Compared with monotherapy, participants who applied once-daily combination treatment (calcipotriol/betamethasone dipropionate) for 8 weeks followed by once-daily calcipotriol for 4 weeks did not experience a significantly greater improvement (SMD -0.12; 95% CI -0.29 to 0.04). Similarly, the psoriasis of participants on monotherapy with twicedaily calcipotriol did not improve significantly more or less than the disease in participants who applied once-daily combination treatment (calcipotriol/betamethasone dipropionate) for 4 weeks followed by 8 weeks using once-daily calcipotriol (weekdays) and once-daily combination therapy (weekends) (SMD 0.13; 95% CI -0.04 to 0.29). These findings were based on the investigators' assessment (IAGI), which were consistent with those of the participants (Analysis 13.4) and with the PASI assessment (Analysis 13.3).

Similarly to Kragballe 2004, Saraceno 2007 compared 12 weeks of calcipotriol monotherapy with 4 weeks of combined therapy followed by 8 weeks of twice-daily calcipotriol. The SMD for the combined end point (0.29; 95% CI -0.04 to 0.62) showed a nonsignificant trend in favour of the complex regimen. Yang 2009 also explored how monotherapy with calcipotriol compared with combined vitamin D/corticosteroid treatment. A 6-week course of monotherapy with calcipotriol was not significantly different to a sequential regimen involving halometasone (SMD 0.41; 95% CI -0.05 to 0.86). Lahfa 2003 found that the effect of clobetasol propionate combined with calcipotriol was similar to the effect of a regimen where the same corticosteroid was used in conjunction with calcitriol: SMD -0.19 (95% CI -0.54 to 0.16).

Analysis 14: Vitamin D alone or in combination versus other treatment: long-term studies (> 24 weeks)

This comparison included active-controlled studies of psoriasis of the body that were at least 24 weeks in duration (see Analysis 14.5 and Table 17). One longer-term placebo-controlled trial of the body (Katz 1991a) and two long-term scalp trials (Luger 2008; Poulin 2010) did not meet the inclusion criteria for this comparison, so we evaluated them elsewhere (Analyses 2, 19, and 18, respectively).

This comparison included one trial, Kragballe 2006, which was a between-patient trial that compared 3 52-week regimens with all treatments used once daily:

• first, combination treatment with calcipotriol and betamethasone dipropionate;

• second, alternating treatment with combination therapy for 4 weeks, then calcipotriol for 4 weeks; and

• third, combination therapy for 4 weeks, then calcipotriol for 48 weeks.

In total, 297 participants contributed data to the analysis. The only outcome measure from the trial that could be included in our review was the Investigator's Global Assessment of Disease Severity (scored from 0, absent, to 5, severe); therefore, results from this section are identical to those in Analysis 14.1. Data were unsuitable for pooling because the same participants contributed to more than one analysis. The adequacy of the concealment of treatment allocation was unclear in this trial.

Data from the combined end point showed there was no significant difference between the three long-term regimens (Figure 5). One year's combination therapy was not significantly better than either the alternating regimen (SMD -0.09; 95% CI -0.36 to 0.18) or the regimen of treatment with 4 weeks of combination therapy followed by 48 weeks of calcipotriol (SMD -0.18; 95% CI -0.47 to 0.10). In both these comparisons, combination therapy achieved a larger absolute benefit, but the difference was not statistically significant. When we compared the alternating therapy with the regimen of 4 weeks' combination therapy followed by 48 weeks of calcipotriol, the SMD for the IAGI also indicated the 2 regimens were not statistically significantly different: -0.09 (95% CI -0.37 to 0.19).

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Figure 5. Forest plot of comparison: 14 Vitamin D alone or in combination vs. other treatment: long term studies (>24wks), outcome: 14.5 Combined end point (IAGI/TSS/PASI/PAGI).



Findings were based on unpublished data supplied by the trial sponsor. Although these data were described as 'intention-to treat', just 47% of the enrolled participants contributed data to the 52-week assessment. The reasons why participants withdrew are summarised in Analysis 14.6, Analysis 14.7, and Analysis 14.8.

Analysis 15: Vitamin D analogues versus other treatment

This comparison incorporated all other vitamin D head-to-head comparisons of treatments for psoriasis of the body (excluding inverse psoriasis) that we had not already included (see Analysis 15.5 and Table 18). We included 12 intervention-comparator contrasts, with data using the combined end point available for 11 of these contrasts. No effectiveness data were available for the comparison of calcipotriol and combined treatment with tazarotene gel plus mometasone furoate cream (Guenther 2000), but the trial did report data on withdrawal and adverse events. Thirteen betweenpatient trials and 6 within-patient trials provided data for the combined end point from 2364 participants. Trial duration ranged between 4 and 12 weeks. Three studies adequately concealed treatment allocation. In light of the pharmacological diversity of the comparators, we only pooled data within subgroups.

In three intervention-comparator contrasts, vitamin D was significantly more effective than the comparator. Specifically, twice-daily calcipotriol was more effective than coal tar polytherapy (SMD -0.59; 95% CI -0.87 to -0.31; I² statistic = 0%; 2 studies) and propylthiouracil cream (SMD -2.24; 95% CI -3.23 to -1.25; I² statistic = NA). On a 6-point IAGI scale, these equate to improvements of 0.65 of a point and 2.46 points, respectively. Twice-daily vitamin D was significantly more effective than once-daily treatment (SMD -0.20; 95% CI -0.32 to -0.07; I² statistic = 0%; 3 studies), achieving an additional improvement of around 0.22 of a point on a 6-point IAGI scale. This finding was based on dosing assessments of 2 vitamin D products: calcipotriol (SMD -0.19; 95% CI -0.37 to -0.02; I² statistic = 12%; 2 studies) and combination treatment with calcipotriol and betamethasone dipropionate (SMD -0.20; 95% CI -0.41 to 0.00; I² statistic = NA).

In the remaining eight intervention-comparator contrasts, we found no significant difference between twice-daily calcipotriol and the comparators. This finding held for the comparison against the following products: coal tar monotherapy (SMD -0.53; 95%)

CI -1.74 to 0.68; I² statistic = 91.1%; 3 studies); nicotinamide, either alone (SMD -0.09; 95% CI -0.49 to 0.31; I² statistic: NA) or in combination with calcipotriol (SMD 0.19; 95% CI -0.14 to 0.52; I² statistic = NA); betamethasone dipropionate ointment and salicylic acid (SMD -0.05; 95% CI -0.26 to 0.15; I² statistic = NA); tacrolimus ointment (SMD -0.55; 95% CI -1.28 to 0.17; I² statistic = 75.9%; 2 studies); tazarotene (SMD -0.10; 95% CI -0.35 to 0.16; I² statistic = 0%; 2 studies); and vitamin B12 cream (SMD -0.55; 95% CI -1.33 to 0.24; I² statistic = NA). The addition of occlusion to calcipotriol did not significantly improve the drug's effectiveness (SMD -0.18; 95% CI -2.04 to 1.68; I² statistic = 96.2%; 2 studies).

Two of the three trials comparing calcipotriol with coal tar monotherapy found a significant difference in favour of calcipotriol (De Simone 1993; Tham 1994); the other trial found a significant difference in favour of coal tar (Alora-Palli 2010). This apparent contradiction may reflect different formulations of coal tar, treatment durations, or different baseline disease severity. Alora-Palli 2010 also assessed effectiveness and quality of life over the six-week post-treatment period, finding that the coal tar group maintained their improvement significantly better than those treated with calcipotriol (see section (c) Quality of life measures).

The two occlusion trials used very different regimens and may be better considered separately. Hindsén 2006 (H) compared calcipotriol with occlusion (applied once weekly for 2 weeks) followed by 4 weeks off treatment against 6 weeks of calcipotriol monotherapy. This trial does not provide a simple assessment of occlusion, since, additionally, treatment is withdrawn. This parallel-group trial contributed data from 209 participants with moderately severe disease and found monotherapy was significantly more effective at the 6-week end point (SMD -1.11; 95% CI -1.40 to -0.81). In the other trial, 19 participants were treated for 8 weeks with twice-daily calcipotriol on both sides of the body, and one side was randomised to receive overnight occlusion with polythene. Bourke 1993b found a significant difference in favour of occlusion (SMD 0.79; 95% CI 0.13 to 1.45); the trial did not explicitly state the severity of participants.

We reported elsewhere trials comparing vitamin D and other

treatments for scalp psoriasis (Analysis 19.5) and inverse psoriasis (Analysis 17.5).

Analysis 16: Flexural/facial psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of topical treatments for inverse or facial psoriasis (see Analysis 16.5 and Table 19). We found evidence on four treatments in this comparison: the potent steroid betamethasone valerate; the vitamin D analogue calcipotriol; and two topical calcineurin inhibitors, pimecrolimus and tacrolimus. We identified only one placebo-controlled trial evaluating tacrolimus ointment (Lebwohl 2004), but the study did not report any effectiveness data suitable for this review. However, the study did contribute data on adverse events and withdrawal rates. We pooled only subtotals in this comparison.

Using the combined end point, data were available for three of the four topical treatments for inverse psoriasis. Two between-patient trials contributed data from 122 participants. Gribetz 2004 was a eight-week study that evaluated twice-daily pimecrolimus cream. The four-week trial by Kreuter 2006 (P) compared betamethasone valerate, calcipotriol, and pimecrolimus against placebo; all treatments were applied once daily. Treatment allocation was adequately concealed in one trial (Gribetz 2004). Therefore, we based the findings for each treatment on a single study.

The SMD for the combined end point found a statistically significant difference in favour of betamethasone valerate (SMD -2.83; 95% CI -3.79 to -1.88) and calcipotriol ointment (SMD -1.08; 95% CI -1.77 to -0.40). These equate to improvements of 3.25 points and 1.24 points on a 6-point IAGI scale. Pimecrolimus cream was also significantly more effective than vehicle (SMD -0.86; 95% CI -1.30 to -0.41; I² statistic = 0%), equivalent to almost 1 point on a 6-point IAGI. This result pooled data on twice-daily applications for 8 weeks (SMD -1.07; 95% CI -1.69 to -0.45) and once-daily pimecrolimus cream for 4 weeks (SMD -0.62; 95% CI -1.27 to 0.02). These findings suggest that different dosing schedules may influence efficacy.

Analysis 17: Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for inverse psoriasis, where we compared vitamin D with an active control (see Analysis 17.5 and Table 20). We identified five intervention-comparator contrasts. Four treatments were compared with calcipotriol: once-daily betamethasone valerate, combined treatment with calcipotriol and hydrocortisone, calcitriol, and pime-crolimus. Calcitriol was compared with tacrolimus.

Using the combined end point, data were available for all five intervention-comparator contrasts. Four trials, varying in duration from 4 to 8 weeks, contributed data from 588 participants. Three studies were between-patient trials, and one was a within-patient study (Ortonne 2003). The adequacy of concealment of treatment allocation was unclear in all four trials.

When applied to sensitive areas of the body, calcipotriol was significantly less effective than three of the comparators. These included betamethasone valerate (SMD 2.02; 95% CI 1.20 to 2.84) (Kreuter 2006 (H)), combination treatment with calcipotriol and hydrocortisone (SMD 0.30; 95% CI 0.11 to 0.50) (Ortonne 2010), and calcitriol (SMD 0.61; 95% CI 0.28 to 0.94) (Ortonne 2003). On a 6-point IAGI scale, the additional benefit equates to

2.22 points for betamethasone valerate, $\frac{1}{3}$ of a point for combi-

nation treatment with hydrocortisone, and $\frac{3}{3}$ of a point for calcitriol.

There was no significant difference between vitamin D and the topical calcineurin inhibitors (calcipotriol versus pimecrolimus: SMD -0.53 (95% CI -1.17 to 0.11; I² statistic = NA); calcitriol versus tacrolimus: SMD 0.42 (95% CI -0.15 to 0.98; I² statistic = NA)).

Analysis 18: Scalp psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of treatments for scalp psoriasis (see Analysis 18.5, Figure 6, and Table 21). We included evidence on 11 treatments in this comparison, with data from the combined end point available for all 11 treatments. Thirteen between-patient trials and 1 within-patient trial (Lepaw 1978) contributed data from 3011 participants. Trial duration ranged between two and eight weeks. Concealment of treatment allocation was adequate in one trial.

Figure 6. Forest plot of comparison: 18 Scalp psoriasis: placebo-controlled trials, outcome: 18.5 Combined end point (IAGI/TSS/PASI/PAGI).

	Difference dom, 95% Cl	Std. Mean Difference IV, Random, 95% Cl
18.5.1 Vitamin D: calcipotriol		
	[-1.69, -0.48]	
	[-0.71, -0.29]	
	[-1.28, -0.16]	•
feterogeneity: Tau* = 0.12; Cni* est for overall effect: Z = 2.53 (P	= 3.25, df = 1 (P = 0.07); l ² = 69% P = 0.01)	
8.5.2 Potent steroid: betameth		
	i [-1.79, 0.06] [-1.30, -0.91]	
	[-1.29, -0.90]	▼
leterogeneity: Tau² = 0.00; Chi²	= 0.25, df = 1 (P = 0.62); I ² = 0%	
est for overall effect: Z = 11.15 ((P < 0.00001)	
8.5.3 Potent steroid: betameth	asone valerate	
ranz 1999 -1.40	[-1.75, -1.05]	
ubtotal (95% Cl) -1.40	[-1.75, -1.05]	•
leterogeneity: Not applicable		
est for overall effect: Z = 7.81 (F	< 0.00001)	
8.5.4 Very potent steroid: amc		_
	[-1.80, -1.04]	7
leterogeneity: Not applicable	[-1.80, -1.04]	•
est for overall effect: Z = 7.27 (P	^o < 0.00001)	
8.5.5 Very potent steroid: clob	etasol propionate	
	[-2.53, -1.45]	
	[-2.55, -1.45] [-1.78, -1.11]	+
	[-1.69, -0.93]	+
	[-1.90, -1.43]	
	[-1.81, -1.34] = 6 20 4f = 2 /P = 0 16\: I8 = 420	•
Heterogeneity: Tauf = 0.02; Chif Test for overall effect: Z = 13.25 (= 5.30, df = 3 (P = 0.15); I ^z = 43% (P < 0.00001)	
10 F C) /		
I8.5.6 Very potent steroid: halc _epaw 1978 -1.11	(-1.69, -0.53)	_
	[-1.69, -0.53]	
Heterogeneity: Not applicable		
Fest for overall effect: Z = 3.78 (P	P = 0.0002)	
18.5.7 Vitamin D in combination	: calcipotriol + BMD	
	[-1.48, -1.08]	•
	[-0.98, -0.27]	*
	[-1.61, -0.32] = 10.15, df = 1 /P = 0.001\; IZ = 0.0%	-
Feterogeneity: Tau* = 0.20; Cni* Fest for overall effect: Z = 2.95 (F	= 10.15, df = 1 (P = 0.001); I ² = 90% ? = 0.003)	
	thasone-17,21-dipropionate plus salicylic acid [-2.50, -0.47]	
	[-2.50, -0.47]	
Heterogeneity: Not applicable		
est for overall effect: Z = 2.86 (P	P = 0.004)	
8.5.9 Other treatment: ciclopir	ox olamine shampoo	
Shuttleworth 1998 -0.07	[-0.82, 0.68]	
	[-0.82, 0.68]	•
Heterogeneity: Not applicable Fest for overall effect: Z = 0.19 (P	2 = 0.85)	
	nolone acetonide, plus occlusion	_
	[-1.69, -0.76] [- 1.69, -0.76]	
leterogeneity: Not applicable	[100, 310]	▼
est for overall effect: Z = 5.12 (F	P < 0.00001)	
18.5.11 Other treatment: salicy		
-	i [-1.79, 0.06]	_
	[-1.79, 0.06]	
Heterogeneity: Not applicable		
Fest for overall effect: Z = 1.82 (F	P = 0.07)	
		<u>ı</u> <u>ı</u> <u>ı</u> <u>ı</u> <u>ı</u>
		-4 -2 Ó 2 4 Favours active treatment Favours placebo
Fest for subgroup differences: C	hi² = 26.09, df = 10 (P = 0.004), l² = 61.7%	, avours acuve reactiont. Favours placebo

Two treatments were not significantly more effective than placebo. These were ciclopirox olamine shampoo (SMD -0.07; 95% CI -0.82 to 0.68; I² statistic = NA) and salicylic acid (SMD -0.86; 95% CI -1.79 to 0.06; I² statistic = NA). Of the nine treatments demonstrated to be significantly more effective than placebo, corticosteroids achieved the largest effects. Two very potent corticosteroids, amcinonide (SMD -1.42; 95% CI -1.80 to -1.04; I² statistic = NA) and clobetasol propionate (SMD -1.57; 95% CI -1.81 to -1.34; I² statistic = 43.3%; 4 studies), delivered benefits equating to almost 2 points (1.70 and 1.88 points, respectively) on a 6-point IAGI (Investigator's Assessment of Global Improvement) scale. Betamethasone dipropionate combined with salicylic acid achieved a similar effect (SMD -1.48; 95% CI -2.50 to -0.47; I² statistic = NA), translating into 1.77 point improvement on the 6-point IAGI over and above placebo.

When used as monotherapy, the 2 potent corticosteroids evaluated were both more effective than placebo: Betamethasone dipropionate (SMD -1.09; 95% CI -1.29 to -0.90; I² statistic = 0%; 2 studies) had a smaller effect than betamethasone valerate (SMD -1.40; 95% CI -1.75 to -1.05; I² statistic = NA), although the difference was not statistically significant. Calcipotriol had the smallest observed benefit (SMD -0.72; 95% CI -1.28 to -0.16; I² statistic = 69.2%; 2 studies). When calcipotriol was combined with betamethasone dipropionate, the pooled effect was larger than for calcipotriol used alone (SMD -0.97; 95% CI -1.61 to -0.32; I² statistic = 90.2%; 2 studies), or 1.16 points on a 6-point improvement scale. Figure 6 shows that both trials found combination therapy to be significantly more effective, but Jemec 2008 (P) found a significantly larger effect (SMD -1.28; 95% CI -1.48 to -1.08) than Tyring 2010 (SMD -0.62; 95% CI -0.98 to -0.27). Participants in the two trials contributing data to the analysis of combination therapy differed in their ethnicity: Tyring 2010 recruited 177 people of Hispanic, Latino, Black, and African American ethnicity; in the trial by Jemec 2008 (P), over 96% of the 1505 enrollees were white. In addition, the trial by Jemec 2008 (P) included a larger proportion of people with severe or very severe disease (37% versus 20%). These factors may help explain the considerable level of heterogeneity (Higgins 2011) observed in the pooled effect for combination therapy.

Sensitivity analysis

We pooled data for the two potent corticosteroids, betamethasone dipropionate and betamethasone valerate. The SMD for the combined end point was -1.18 (95% CI -1.40 to -0.96; I² statistic = 19.9%; 3 studies), equating to 1.41 points on a 6-point IAGI scale. For the 3 very potent corticosteroids, the SMD was -1.51 (95% CI -1.70 to -1.31; I² statistic = 37.5%; 6 studies), translating as a 1.80 point improvement over placebo on the IAGI.

Analysis 19: Scalp psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for scalp psoriasis in which one of the interventions was a vitamin D product (used either as monotherapy or in combination with another product) (see Analysis 19.5 and Table 22).

We identified six intervention-comparator contrasts, and data on the combined end point were available for all these contrasts. All studies were parallel-group in design (between-patient). Twelve studies contributed data from 5413 participants. Trial duration ranged from 4 to 8 weeks in 11 studies, but duration was 52 weeks in one trial (Luger 2008). The adequacy of the concealment of treatment allocation was unclear in all 12 trials.

Based on the combined end point scores, monotherapy with vitamin D was consistently less effective than monotherapy with potent or very potent corticosteroids, and it was also significantly less effective than combination therapy with vitamin D and a potent steroid. Specifically, calcipotriol was significantly less effective than betamethasone dipropionate (SMD 0.48; 95% CI 0.32 to 0.64; I² statistic = 60.4%; 2 studies), betamethasone valerate (SMD 0.37; 95% CI 0.20 to 0.55; I² statistic = 0%; 2 studies), and clobetasol propionate (SMD 0.37; 95% CI 0.05 to 0.69; I² statistic = NA). On a 6-point investigators' global assessment of improvement (IAGI) scale, these SMDs translate into 0.62, 0.48, and 0.48 points, respectively.

Combination treatment achieved a significantly greater benefit than either calcipotriol alone or betamethasone dipropionate alone. Calcipotriol was significantly less effective than combination treatment (SMD 0.64; 95% CI 0.44 to 0.84; I² statistic = 82.3%; 4 studies), with combined therapy delivering a benefit equivalent to 0.83 of a point on a 6-point IAGI. Combination treatment was significantly more effective than monotherapy with betamethasone dipropionate (SMD -0.18; 95% CI -0.26 to -0.10; I² statistic = 0%; 3 studies), translating into a net benefit of 0.24 of a point on a 6-point IAGI scale.

Compared with coal tar polytherapy, calcipotriol achieved a larger benefit, though this was not statistically significant at the 5% level (SMD -0.45; 95% CI -0.92 to 0.02; I² statistic = 89.8%; 3 studies).

(2) Secondary outcome measures

(a) Withdrawal rates (total rate; withdrawal due to adverse events; withdrawal due to treatment failure)

Analysis 1: Vitamin D analogues versus placebo

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We pooled withdrawal data from seven of the eight vitamin D analogues using a random-effects risk difference (RD) metric (see Analysis 1.6, Analysis 1.7, and Analysis 1.8). There were no withdrawal data for the comparison of calcipotriol plus occlusion versus placebo.

There was no significant difference in the pooled rate of withdrawals from the trials for any reason (total withdrawals) (RD: -0.02; (95% CI -0.05 to 0.00; I² statistic = 51.4%). Rates of withdrawals due to adverse events were not statistically significantly different (RD -0.00; 95% CI -0.02 to 0.01; I² statistic = 36.4%), and neither were withdrawals due to treatment failure (RD -0.03; 95% CI -0.05 to 0.00; I² statistic = 81.7%). Individually, none of the eight vitamin D analogues were significantly different to placebo in any of the three withdrawal rates.

Analysis 2: Corticosteroid (potent) versus placebo

This comparison included 10 potent corticosteroids. No withdrawal data were available for difforasone diacetate, and some withdrawal data were missing for fluticasone propionate, mometasone furoate, and betamethasone dipropionate. Where available, we pooled data using a random-effects risk difference metric (see Analysis 2.6, Analysis 2.7, and Analysis 2.8). There was a small but statistically significant difference in favour of potent corticosteroids for withdrawals from the trials for any reason (total withdrawals): RD -0.14 (-0.22 to -0.05; I² statistic = 81.5%). This finding was driven by two large trials of once-daily betamethasone dipropionate (Fleming 2010 (P); Kaufmann 2002 (P)) and two small trials of betamethasone dipropionate used as maintenance (weekend pulse therapy) (Katz 1987a; Katz 1991a). In the other corticosteroids, the RD showed no statistically significant difference.

There was no significant difference in the pooled rate of withdrawals due to adverse events (RD -0.01; 95% CI -0.05 to 0.02; I² statistic = 60.9%). However, betamethasone dipropionate once daily was associated with statistically significantly lower rates of withdrawal due to adverse events than placebo (RD -0.07; 95% CI -0.11 to -0.02; I² statistic = NA).

For the rate of withdrawals due to treatment failure (Analysis 2.8), findings differed by treatment duration; therefore, we only pooled subgroup data. For short-term treatments, there was no difference relative to placebo (RD 0.00; 95% CI -0.02 to 0.02; I² statistic = 0.0%). This is the pooled effect for the placebo comparisons with betamethasone valerate, budesonide, desonide, and hydrocortisone buteprate (data on withdrawals due to treatment failure for once-daily and twice-daily betamethasone dipropionate were not available). For maintenance treatment with betamethasone dipropionate, there was a statistically significant difference in favour of maintenance treatment when compared with placebo: RD -0.46; 95% CI -0.61 to -0.31; I² statistic = 0.0% (Katz 1987a; Katz 1991a).

Analysis 3: Corticosteroid (very potent) versus placebo

We identified withdrawal data for two of the three very potent corticosteroids in this comparison and pooled data using a randomeffects risk difference metric (see Analysis 3.6, Analysis 3.7, and Analysis 3.8). There were no withdrawal data available for halcinonide. There were no statistically significant differences between very potent corticosteroids and placebo for any type of withdrawal or for any individual treatment. For total withdrawals, the risk difference was -0.05 (95% CI -0.10 to 0.01; I² statistic = 83.7%). Differences in withdrawals due to adverse events (RD -0.00; 95% CI -0.01 to 0.01; I² statistic = 0.0%) and withdrawals due to treatment failure (RD -0.00; 95% CI -0.02 to 0.01); I² statistic = 13.5%) were also small and non-significant.

Analysis 4: Dithranol versus placebo

We pooled withdrawal data on dithranol versus placebo using a random-effects risk difference metric (see Analysis 4.6, Analysis 4.7, and Analysis 4.8). There were no statistically significant differences between dithranol and placebo for any type of withdrawal. For total withdrawals, the risk difference was 0.00 (95% CI -0.09 to 0.09; I² statistic = 0%). Differences in withdrawals due to adverse events (RD 0.00; 95% CI -0.05 to 0.05; I² statistic = 0%) and withdrawals due to treatment failure (RD 0.00; 95% CI -0.11 to 0.11; I² statistic = 0%) were also small and non-significant.

Analysis 5: Vitamin D combination products versus placebo

We pooled data from trials of vitamin D combination products using a random-effects risk difference (RD) metric (see Analysis 5.6, Analysis 5.7, and Analysis 5.8). In all three withdrawal measures, combined treatment - whether used once or twice daily was significantly less likely to lead to withdrawal from the trial. This finding held for total withdrawals (RD -0.12; 95% CI -0.17 to -0.07; I² statistic = 59.3%), withdrawals due to adverse events (RD -0.07; 95% CI -0.11 to -0.04; I² statistic = 55.0%), and withdrawals due to treatment failure (RD -0.09; 95% CI -0.12 to -0.06; I² statistic = 0%).

Analysis 6: Other treatment versus placebo

We report findings separately for 22 of the 26 treatments considered; no withdrawal data were available for four placebo comparisons: those involving a herbal skin care product, topical sirolimus, topical tacrolimus, or coal tar. Data on total withdrawals were available for 22 treatments; for withdrawals due to adverse events or treatment failure, data were available for 18 treatments (see Analysis 6.6, Analysis 6.7, and Analysis 6.8).

We assessed withdrawal data using a random-effects risk difference metric. Relative to placebo, tazarotene was associated with a significantly higher rate of withdrawal due to adverse events (RD 0.07; 95% CI 0.05 to 0.10) and *Mahonia aquifolium* was associated with a significantly lower rate of total withdrawal (i.e. withdrawal from the trial for any reason) (RD -0.23; 95% CI -0.32 to -0.14). No other treatment was significantly different from placebo on any of the three withdrawal measures assessed.

Aloe vera extract 0.5% hydrophilic cream, three times per day

Sixty participants contributed data (Syed 1996). The comparison with placebo found no statistically significant difference for aloe vera extract, for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Anti-IL-8 monoclonal antibody cream

Ninety-six participants contributed data (Jin 2001). For total withdrawals, the comparison with placebo found no statistically significant difference for anti-IL-8 monoclonal antibody cream. The study did not report data on withdrawals due to adverse events or treatment failure.

Betamethasone 17-valerate 21-acetate plus tretinoin plus salicylic acid

Eighty-five participants contributed data (Santoianni 2001). The comparison with placebo found no statistically significant difference for betamethasone 17-valerate 21-acetate plus tretinoin plus salicylic acid for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Caffeine (topical) 10%, three times per day

Thirty-nine participants contributed data (Vali 2005). The comparison with placebo found no statistically significant difference for topical caffeine for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Calcipotriene 0.005% ointment + nicotinamide 0.05% or 0.1% or 0.7% or 1.4%, twice daily

One hundred and sixty participants contributed data (Levine 2010 (P)). The comparison with placebo found no statistically significant difference for combination therapy with calcipotriol and nicotinamide for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Dead Sea salts emollient lotion

Twenty-four participants contributed data (Cheesbrough 1992). The comparison with placebo found no statistically significant difference for Dead Sea salts emollient lotion for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Fish oil plus occlusion

Twenty-five participants contributed data (Escobar 1992). The comparison with placebo found no statistically significant difference for fish oil plus occlusion for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Herbal skin care (Dr Michaels® cleansing gel, ointment, and skin conditioner), twice daily

No withdrawal data were available for this comparison.

Hexafluoro-1,25-dibydroxyvitamin D3

Fifteen participants contributed data (Durakovic 2001). The comparison with placebo found no statistically significant difference for hexafluoro-1,25-dihydroxyvitamin D3 for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Indigo naturalise 1.4% ointment

Fifty-six participants contributed data from two within-patient trials (Lin 2007; Lin 2008). The comparison with placebo found no statistically significant difference for indigo naturalise ointment for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Kukui nut oil, three times per day

Thirty participants contributed data (Brown 2005). The comparison with placebo found no statistically significant difference for kukui nut oil for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Mahonia aquifolium (Reliéva™), twice daily

Two hundred participants contributed data (Bernstein 2006). For total withdrawals, the comparison with placebo found a statistically significant difference for *Mahonia aquifolium* (RD -0.23;

95% CI -0.32 to -0.14). The trial did not report data on withdrawals due to adverse events or treatment failure.

Methotrexate gel

Sixty participants contributed data (Syed 2001b). The study by Sutton 2001 on methotrexate gel did not report withdrawal data. The comparison with placebo found no statistically significant difference for methotrexate gel for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Mycophenolic acid ointment

Seven participants contributed data (Geilen 2000). The comparison with placebo found no statistically significant difference for mycophenolic acid ointment for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

NG-monomethyl-L-arginine (L-NMMA) cream

Seventeen participants contributed data (Ormerod 2000). The comparison with placebo found no statistically significant difference for NG-monomethyl-L-arginine (L-NMMA) cream for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Nicotinamide 1.4%, twice daily

Eighty-eight participants contributed data (Levine 2010 (P)). The comparison with placebo found no statistically significant difference for nicotinamide for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Oleum horwathiensis

Fifty participants contributed data (Lassus 1991). The comparison with placebo found no statistically significant difference for oleum horwathiensis for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Omega-3 polyunsaturated fatty acids ointment

Seventy-three participants contributed data (Henneicke-v. Z. 1993). The comparison with placebo found no statistically significant difference for omega-3 polyunsaturated fatty acids ointment for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

trial did not report data on withdrawals due to adverse events or treatment failure.

Polymyxin B cream, 200,000 U/g

Fifteen participants contributed data (Stutz 1996). The comparison with placebo found no statistically significant difference for polymyxin B cream for total withdrawals. The trial did not report data on withdrawals due to adverse events or treatment failure.

Platelet aggregation activating factor (PAF) (Ro 24-0238)

Fifty-two participants contributed data (Wolska 1995). The comparison with placebo found no statistically significant difference for platelet aggregation activating factor for total withdrawals. The

PTH (1-34) in Novasome A® liposomal cream, twice daily

Fifteen participants contributed data (Holick 2003). The comparison with placebo found no statistically significant difference for PTH (1-34) in Novasome A® liposomal cream for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Sirolimus (topical)

No withdrawal data were available for this comparison.

Tacrolimus ointment

No withdrawal data were available for this comparison.

Tar

No withdrawal data were available for this comparison.

Tazarotene

Two studies, with 1627 participants, contributed data (Weinstein 1996; Weinstein 2003). The comparison with placebo found no statistically significant difference for tazarotene for total withdrawals or withdrawals due to treatment failure. However, tazarotene was significantly more likely to lead to withdrawal from the trial due to adverse events (RD 0.07; 95% CI 0.05 to 0.10; I² statistic = 0%).

Theophylline 1% ointment, twice daily

Twenty-two participants contributed data (Papakostantinou 2005). The comparison with placebo found no statistically significant difference for theophylline ointment for total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Analysis 7: Vitamin D analogues versus corticosteroid (potent)

Seven of the eight vitamin D analogues that were compared with potent corticosteroids reported withdrawal data (withdrawal data on calcipotriol versus desoxymetasone were not available) (see Analysis 7.6, Analysis 7.7, and Analysis 7.8). We pooled withdrawal data on these seven treatment comparison pairs using a random-effects risk difference (RD) metric.

Relative to potent corticosteroid, there was a statistically significant difference in total withdrawals in favour of corticosteroids: RD 0.02 (95% CI 0.00 to 0.03; I² statistic = 0%). Pooled rates of withdrawals due to adverse events or treatment failure were not statistically significantly different. Regarding individual vitamin D analogues, the only statistically significant differences in withdrawals relative to corticosteroid were for the comparison of calcipotriol against betamethasone dipropionate (total withdrawals: RD 0.03 (95% CI 0.01 to 0.06; I² statistic = 0%); withdrawals due to adverse events: RD 0.02 (95% CI 0.00 to 0.04; I² statistic = NA)).

Analysis 8: Vitamin D analogues versus corticosteroid (very potent)

We pooled withdrawal data on calcipotriol against clobetasol propionate using a random-effects risk difference (RD) metric (see Analysis 8.6, Analysis 8.7, and Analysis 8.8). Relative to the very potent corticosteroid, we found no statistically significant difference for calcipotriol on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Analysis 9: Vitamin D combined with corticosteroid versus corticosteroid

We pooled withdrawal data on combination treatment (vitamin D and a corticosteroid) against monotherapy with a corticosteroid using a random-effects risk difference (RD) metric (see Analysis 9.6, Analysis 9.7, and Analysis 9.8).

Relative to the corticosteroid, we found no statistically significant difference for combination treatment on total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure. There were no data on the rate of withdrawals due to treatment failure for the comparison of combined treatment with calcipotriol and betamethasone dipropionate versus betamethasone dipropionate monotherapy.

Analysis 10: vitamin D alone or in combination versus dithranol

Withdrawal data were available for three intervention-comparator pairs: calcipotriol versus dithranol, calcitriol versus dithranol, and tacalcitol versus dithranol. We pooled these data using a randomeffects risk difference (RD) metric (see Analysis 10.6, Analysis 10.7, and Analysis 10.8). For the comparison of tacalcitol versus dithranol, there were no data available on withdrawals due to adverse events or treatment failure.

There was no statistically significant difference for total withdrawals, either for individual intervention-comparator pairs or for the pooled effect (RD -0.02; 95% CI -0.06 to 0.01; I² statistic = 0%). The pooled risk difference showed a significant difference in favour of vitamin D for withdrawals due to adverse events (RD -0.03; 95% CI -0.06 to -0.00; I² statistic = 26.3%), but no significant difference in the rate of withdrawals due to treatment failure (RD -0.00; 95% CI -0.02 to 0.02; I² statistic = 0%).

Analysis 11: Vitamin D alone or in combination versus other vitamin D analogue

Two of the three vitamin D analogues head-to-head comparisons reported withdrawal data (withdrawal data on calcipotriol versus tacalcitol were not available). We pooled withdrawal data on calcipotriol versus calcitriol, and calcipotriol versus maxacalcitol, using a random-effects risk difference (RD) metric (see Analysis 11.6, Analysis 11.7, and Analysis 11.8). We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure. There was also no significant difference in the withdrawal rates for the individual contrasts, calcipotriol versus calcitriol, and calcipotriol versus maxacalcitol.

Analysis 12: Vitamin D alone or in combination versus vitamin D + corticosteroid

To simplify the analysis, we summarised withdrawal data by grouping the first 10 intervention-comparator pairs in a single category: calcipotriol versus calcipotriol and corticosteroid. The remaining comparisons were calcitriol versus calcitriol and corticosteroid, and tacalcitol versus calcipotriol and corticosteroid. We pooled withdrawal data using a random-effects risk difference (RD) metric (see Analysis 12.6, Analysis 12.7, and Analysis 12.8). There were no data on withdrawals due to treatment failure for the comparisons of combined therapy versus calcitriol or combined therapy versus tacalcitol.

In the comparison of calcipotriol against calcipotriol plus corticosteroid, total withdrawals were statistically significantly different in favour of polytherapy: RD 0.03 (95% CI 0.01 to 0.05; I² statistic = 13.3%). We based this on findings from 4922 participants in 13 trials. Similarly, a significant difference in favour of polytherapy was evident from the data on withdrawals due to adverse events:

Topical treatments for chronic plaque psoriasis (Review) Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd. RD 0.02 (95% CI 0.01 to 0.03; I² statistic = 32.9%). However, we found no statistically significant difference for withdrawals due to treatment failure (RD 0.01; 95% CI -0.00 to 0.02; I² statistic = 0%). When either calcitriol or tacalcitol was compared with combination therapy, differences in total withdrawals and in withdrawals due to adverse events were not statistically significant.

Analysis 13: Vitamin D alone or in combination versus other treatments: complex regimens

This comparison covers complex regimens, defined here as treatment sequences that do not consist of a simple head-to-head comparison between two active treatments. We summarised withdrawals rates on twelve intervention-comparator contrasts (see Analysis 13.6, Analysis 13.7, and Analysis 13.8). There were data on total withdrawal rates for all 12 contrasts, but data on withdrawals due to adverse events were missing for 3 contrasts, and there were no data on withdrawals due to treatment failure for 6 contrasts.

In the analysis of total withdrawals (Analysis 13.6), there was a significant difference in favour of the complex regimen for four contrasts:

• Participants were significantly more likely to withdraw from the trial after 12 weeks of calcipotriol than participants who received combination therapy (8 weeks) followed by calcipotriol (4 weeks) (RD 0.05; 95% CI 0.00 to 0.10).

• Trial withdrawal rates were also significantly higher in participants receiving 12 weeks of monotherapy with calcipotriol than those treated with combination therapy (4 weeks) followed by alternating calcipotriol on weekdays and combination therapy at the weekend (8 weeks) (RD 0.08; 95% CI 0.03 to 0.13).

• After an initial 4-week phase with combination ointment, participants who received 8 weeks' maintenance with calcipotriol (RD 0.08; 95% CI 0.03 to 0.14) or 8 weeks' maintenance with an alternating calcipotriol/combined therapy routine (RD 0.11; 95% CI 0.06 to 0.17) were significantly less likely to withdraw than participants who used vehicle ointment for maintenance.

In the analysis of withdrawals due to adverse events (Analysis 13.7), there were no significant differences between vitamin D and any of the complex regimens.

In the analysis of withdrawals due to treatment failure (Analysis 13.8), there were significant differences in two of the six contrasts for which we had reported data. After 12 weeks of calcipotriol, participants were significantly more likely to withdraw from the trial because of treatment failure than those who received combination therapy followed by calcipotriol (RD 0.21; 95% CI 0.10 to 0.33). Withdrawals due to treatment failure were more likely in participants who received 8 weeks' monotherapy with tacalcitol than those who received combination therapy followed by calcipotriol (RD 0.05; 95% CI 0.02 to 0.08).

Analysis 14: Vitamin D alone or in combination versus other treatment: long-term studies (> 24 weeks)

This comparison included active-controlled trials of psoriasis of the body that were at least 24 weeks in duration (see Analysis 14.6, Analysis 14.7, and Analysis 14.8). There were three interventioncomparator contrasts, all 52 weeks in duration and all reported by a single study (Kragballe 2006).

Participants received one of three long-term treatments:

• once-daily combined calcipotriol and betamethasone dipropionate;

• combination therapy (4 weeks) followed by calcipotriol (for 4 weeks), rotated over 52 weeks; or

• combination therapy for 4 weeks followed by calcipotriol for 48 weeks.

There were no significant differences between these three regimens in any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Analysis 15: Vitamin D analogues versus other treatment

This comparison compared vitamin D analogue with 12 other treatments. We assessed the results from the analyses of withdrawal data using a random-effects risk difference (RD) metric, and we report these separately (see Analysis 15.6, Analysis 15.7, and Analysis 15.8). Withdrawal data were missing only in the head-to-head comparison of vitamin D (alone or in combination) versus calcipotriol with occlusion.

Of the remaining 11 intervention-comparator contrasts, we found only 1 statistically significant difference in withdrawal rates: The rate of total withdrawals was significantly lower for calcipotriol than for tacrolimus (RD -0.13; 95% CI -0.25 to -0.01).

Calcipotriol versus coal tar

In the comparison of calcipotriol and coal tar, we found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Calcipotriol versus coal tar polytherapy

In the comparison of calcipotriol and coal tar polytherapy, we found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Topical treatments for chronic plaque psoriasis (Review) Copyright © 2013 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd. We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Calcipotriol versus calcipotriol + nicotinamide (0.05%, 0.1%, 0.7%, or 1.4%), twice daily

We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Calcipotriol versus corticosteroid + salicylic acid

We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure. However, the risk difference for the withdrawal rate for adverse events was almost statistically significant in favour of combined therapy (RD 0.05; 95% CI -0.00 to 0.10).

Calcipotriol versus propylthiouracil cream

We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Calcipotriol versus tacrolimus ointment

We found no statistically significant difference in withdrawals due to adverse events or withdrawals due to treatment failure. However, the rate of total withdrawals was significantly lower for calcipotriol than for tacrolimus (RD -0.13; 95% CI -0.25 to -0.01) (Ortonne 2006).

Calcipotriol versus tazarotene

We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Calcipotriol versus tazarotene gel plus mometasone furoate cream

We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Calcipotriol versus vitamin B12 cream

We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Head-to-head vitamin D alone or in combination: dosing

We found no statistically significant difference on any of the withdrawal assessments: total withdrawals, withdrawals due to adverse events, or withdrawals due to treatment failure.

Head-to-head vitamin D alone or in combination: occlusion

We found no usable withdrawal data reported for this comparison.

Analysis 16: Flexural/facial psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of four topical treatments for inverse or facial psoriasis. Data on three withdrawal rates were available for all four treatments: the potent steroid be-tamethasone valerate; the vitamin D analogue calcipotriol; and two topical calcineurin inhibitors, pimecrolimus and tacrolimus (see Analysis 16.6, Analysis 16.7, and Analysis 16.8).

Relative to placebo, participants receiving topical tacrolimus were significantly less likely to withdraw from treatment for any reason (RD -0.17; 95% CI -0.30 to -0.03) or because of treatment failure (RD -0.11; 95% CI -0.19 to -0.02). No other treatment was significantly different from placebo on any of the three withdrawal measures assessed.

Analysis 17: Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for flexural or facial psoriasis, where we compared vitamin D with an active control (see Analysis 17.6, Analysis 17.7, and Analysis 17.8). We identified five intervention-comparator contrasts, and no withdrawal data were missing. Four treatments were compared with calcipotriol: once-daily betamethasone valerate, combined treatment with calcipotriol and hydrocortisone, calcitriol, and pimecrolimus. Calcitriol was compared with tacrolimus.

The rate of withdrawals due to adverse events was significantly higher in people treated with calcipotriol compared to those receiving combination therapy with calcipotriol and hydrocortisone for facial psoriasis (RD 0.06; 95% CI 0.02 to 0.11) and compared to those receiving calcitriol (RD 0.09; 95% CI 0.01 to 0.18). We found no other significant differences in withdrawal rates.

Analysis 18: Scalp psoriasis: placebo-controlled trial

This comparison included placebo-controlled trials of 11 topical treatments for scalp psoriasis (see Analysis 18.6, Analysis 18.7, and Analysis 18.8). All treatments had data on at least one withdrawal measure, which we assessed using a risk difference (RD) metric. Relative to placebo, participants treated with betamethasone dipropionate were significantly less likely to withdraw from the trial for any reason (total withdrawals) (RD -0.14; 95% CI -0.21 to -0.06), because of adverse events (RD -0.04; 95% CI -0.08 to -0.00), or because of treatment failure (RD -0.10; 95% CI -0.16 to -0.05). Participants receiving combination therapy with calcipotriol and betamethasone dipropionate were significantly less likely to withdraw from the trial for any reason (total withdrawals) (RD -0.09; 95% CI -0.16 to -0.03) or because of treatment failure (RD -0.11; 95% CI -0.17 to -0.06). We found no other significant differences in withdrawal rates.

Analysis 19: Scalp psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for scalp psoriasis in which one of the interventions was a vitamin D product (used either as monotherapy or in combination with another product) (see Analysis 19.6, Analysis 19.7, and Analysis 19.8). We identified six intervention-comparator contrasts. There were no data for withdrawal due to treatment failure for one of these contrasts (calcipotriol versus coal tar polytherapy). No other withdrawal data were missing.

Relative to combination treatment with calcipotriol and betamethasone dipropionate, calcipotriol monotherapy was associated with a significantly higher rate of total withdrawals (RD 0.11; 95% CI 0.05 to 0.18; I² statistic = 78.5%), withdrawals due to adverse events (RD 0.06; 95% CI 0.02 to 0.09; I² statistic = 78.9%) and withdrawals due to treatment failure (RD 0.05; 95% CI 0.01 to 0.10; I² statistic = 88.2%). Compared with betamethasone dipropionate, combination therapy was also significantly less likely to result in participant withdrawal due to treatment failure (RD -0.01; 95% CI -0.02 to -0.00; I² statistic = 0%).

Study participants treated with calcipotriol were significantly more likely to withdraw because of adverse events than participants receiving betamethasone valerate (RD 0.03; 95% CI 0.01 to 0.06; I² statistic = 0%) or coal tar polytherapy (RD 0.08; 95% CI 0.02 to 0.14; I² statistic = NA).

(b) Adverse events (local (cutaneous) and systemic)

(i) Findings from the main review

Analysis 1: Vitamin D analogues versus placebo

We pooled data on adverse events using a random-effects risk difference metric (see Analysis 1.9 and Analysis 1.10). There were no adverse events data for the comparison of calcipotriol plus occlusion versus placebo. Therefore, data were available for seven of the eight vitamin D analogues evaluated against placebo

Of the seven vitamin D analogues evaluated against placebo, we found only one statistically significantly difference. Twice-daily becocalcidiol was significantly more likely to cause local adverse events than placebo (RD 0.10; 95% CI 0.00 to 0.19). There were no significant differences for the pooled risk difference for either local adverse events (RD 0.00; 95% CI -0.01 to 0.02; I² statistic = 0%) or systemic adverse events (RD 0.00; 95% CI -0.01 to 0.01; I² statistic = 0%).

Analysis 2: Corticosteroid (potent) versus placebo

We pooled data on adverse events using a random-effects risk difference metric (see Analysis 2.9 and Analysis 2.10). Data on local adverse events were available for 7 of the 10 corticosteroids; data on systemic adverse events were available only for betamethasone dipropionate twice daily, betamethasone dipropionate (maintenance), and mometasone furoate.

The pooled analysis of local adverse events found a significant difference in favour of corticosteroids (RD -0.04; 95% CI -0.08 to -0.00; I² statistic = 56.2%). Among the individual potent corticosteroids evaluated against placebo, we found only one statistically significantly difference. Once-daily betamethasone dipropionate was less likely than placebo to be associated with local adverse events: RD -0.10 (95% CI -0.15 to -0.04; I² statistic = 2.5%). We found no significant difference in the pooled analyses of systemic adverse events data.

Analysis 3: Corticosteroid (very potent) versus placebo

We pooled data on adverse events using a random-effects risk difference metric (see Analysis 3.9 and Analysis 3.10). The comparison with placebo found no statistically significant difference for any of the three very potent corticosteroids considered for either local adverse events or systemic adverse events.

Analysis 4: Dithranol versus placebo

We pooled data on adverse events using a random-effects risk difference metric (see Analysis 4.9 and Analysis 4.10). The comparison with placebo found no statistically significant difference for pooled findings on local adverse events or systemic adverse events. In one study, a significant difference in favour of placebo was evident: RD 0.40 (95% CI 0.08 to 0.72; I² statistic = NA) (Volden 1992).

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Analysis 5: Vitamin D combination products versus placebo

We pooled data on adverse events using a random-effects risk difference metric (see Analysis 5.9 and Analysis 5.10). The rate of local adverse events was significantly lower for combination therapy compared with placebo (RD -0.05; 95% CI -0.08 to -0.02; I² statistic = 10.0%). There was no significant difference in the rate of systemic adverse events.

Analysis 6: Other treatment versus placebo

Our review reports findings separately for 22 of the 26 comparisons considered (see Analysis 6.9 and Analysis 6.10). No data on adverse events were available for placebo comparisons involving polymyxin B cream, topical sirolimus, topical tacrolimus, coal tar, or tazarotene. Data on local adverse events were available for 22 comparisons, but data on systemic adverse events were reported for just 9 of the 26 comparisons. We assessed data on adverse events using a random-effects risk difference metric. The comparison with placebo found no statistically significant difference in the rate of either local or systemic adverse events for any treatment.

Analysis 7: Vitamin D analogues versus corticosteroid (potent)

Of the eight vitamin D analogues that were compared with potent corticosteroids, data on local adverse events were available for five comparisons, and data on systemic events were available for four comparisons (see Analysis 7.9 and Analysis 7.10). There were no data on adverse events for calcipotriol versus desoxymetasone, calcipotriol versus diflorasone diacetate, and calcitriol versus betamethasone valerate. In addition, systemic adverse events data were unavailable for calcipotriol versus fluocinonide. We pooled data on adverse effects using a random-effects risk difference metric.

In the comparison of individual vitamin D analogues and potent corticosteroids, our review found one statistically significant difference in favour of potent corticosteroids. Pooled data from 3 trials with 1739 participants indicated that the rate of local adverse events was significantly higher in the calcipotriol group than in the group receiving betamethasone dipropionate (RD 0.07; 95% CI 0.04 to 0.09; I² statistic = 0%). This finding drove the pooled (class) result for local adverse events (RD 0.07; 95% CI 0.02 to 0.11; I² statistic = 82.4%). Our review found no other statistically significant differences for local or systemic adverse events in this comparison.

Analysis 8: Vitamin D analogues versus corticosteroid (very potent)

We pooled withdrawal data on adverse effects using a randomeffects risk difference metric (see Analysis 8.9 and Analysis 8.10). There was no statistically significantly difference between calcipotriol and clobetasol propionate, either for local adverse events (RD -0.05; 95% CI -0.18 to 0.08) or systemic events (RD -0.05; 95% CI -0.18 to 0.08).

Analysis 9: Vitamin D combined with corticosteroid versus corticosteroid

We pooled withdrawal data on combination treatment (vitamin D and a corticosteroid) against monotherapy with a corticosteroid using a random-effects risk difference metric (see Analysis 9.9 and Analysis 9.10). No adverse events data were available for the comparison of combination (calcipotriol/clobetasol propionate) therapy and clobetasol propionate.

The pooled analysis found no statistically significant difference in adverse event rates between combination (calcipotriol/betamethasone dipropionate) therapy and betamethasone dipropionate. There was no significant difference in local adverse events for combination (calcipotriol/betamethasone) therapy and clobetasol propionate, and there were no data on systemic adverse events for this comparison.

Analysis 10: Vitamin D alone or in combination versus dithranol

Adverse events data were available for three intervention-comparator pairs: calcipotriol versus dithranol, calcitriol versus dithranol, and tacalcitol versus dithranol. We pooled these data using a random-effects risk difference metric (see Analysis 10.9 and Analysis 10.10).

Vitamin D was statistically significantly less likely to cause local adverse events (RD -0.32; 95% CI -0.43 to -0.20; I² statistic = 84.5%), although substantial heterogeneity was evident in the pooled statistic (Higgins 2011). This finding also held for each of the three intervention-comparator pairs, with the effect size ranging from -0.25 (calcipotriol versus dithranol) to -0.67 (calcitriol versus dithranol). However, the analysis of systemic adverse events found no statistically significant differences.

Analysis 11: Vitamin D alone or in combination versus other vitamin D analogue

We reported local adverse events data on two of the vitamin D analogues head-to-head comparisons (see Analysis 11.9 and Analysis 11.10). Systemic adverse events data were not available for any of the three comparisons. We pooled data on adverse effects using a random-effects risk difference metric. Overall, our review found no statistically significant difference on either local or systemic adverse events when comparing pooled vitamin D analogues headto-head. However, in the comparison of calcipotriol against calcitriol, the rate of local adverse events was significantly lower for calcitriol: RD 0.07 (95% CI 0.01 to 0.14; I² statistic = NA).

Analysis 12: Vitamin D alone or in combination versus vitamin D + corticosteroid

To simplify the comparison, we summarised adverse events data by grouping the first 10 intervention-comparator pairs in a single category: calcipotriol versus calcipotriol and corticosteroid. The remaining comparisons were 1) calcitriol versus calcitriol and 2) corticosteroid and tacalcitol versus calcipotriol and corticosteroid. There were no data on systemic adverse events for the comparison of tacalcitol against calcipotriol and corticosteroid.

We pooled data on adverse effects using a random-effects risk difference (RD) metric (see Analysis 12.9 and Analysis 12.10). In the local adverse events comparison of vitamin D against vitamin D plus corticosteroid, our review found a statistically significantly difference in favour of combination therapy: RD 0.06 (95% CI 0.05 to 0.08; I² statistic = 4.1%). This finding also held for each individual intervention-comparator pair. Our analysis found no statistically significantly difference in the rates of systemic adverse events.

Analysis 13: Vitamin D alone or in combination versus other treatments: complex regimens

This comparison included 12 intervention-comparator contrasts (see Analysis 13.9 and Analysis 13.10). Data on local adverse events were available for 11 contrasts, but there were no data on the comparison of calcipotriol (12 weeks) versus combination therapy (4 weeks) followed by calcipotriol (8 weeks). No trial in this comparison reported data on systemic adverse events.

In 4 of the 11 contrasts for which data were available, the complex regimen was associated with a significantly lower rate of local adverse events than when the vitamin D analogue was used alone. Twelve weeks of calcipotriol monotherapy was associated with significantly higher rates of local adverse events than two complex regimen comparators. Specifically, the rate of local adverse events was higher for calcipotriol monotherapy than for a regimen consisting of 8 weeks of combination therapy followed by 4 weeks of once-daily calcipotriol (RD 0.11; 95% CI 0.06 to 0.17). Calcipotriol monotherapy was also more likely to be associated with local adverse events than 4 weeks of combination therapy followed by 8 weeks of once-daily calcipotriol on weekdays and combination therapy at the weekend (RD 0.11; 95% CI 0.05 to 0.17) (Kragballe 2004). When we compared the two complex regimens directly, the rates of cutaneous adverse events were not significantly different.

Relative to calcipotriol, combination therapy with halometasone and calcipotriol had a significantly lower rate of adverse events of the skin (RD 0.26; 95% CI 0.07 to 0.45). An 8-week course of tacalcitol had significantly higher rates of adverse events than a routine of 4 weeks' combined (calcipotriol/betamethasone dipropionate) therapy followed by 4 weeks' monotherapy with calcipotriol (RD 0.06; 95% CI 0.01 to 0.11). There was no significant difference in the rate of local adverse events in any of the remaining seven intervention-comparator contrasts.

Analysis 14: Vitamin D alone or in combination versus other treatment: long-term studies (> 24 weeks)

This comparison included active-controlled studies of psoriasis of the body that were at least 24 weeks in duration (see Analysis 14.9 and Analysis 14.10). No data on systemic adverse events were reported. We included one trial in this comparison: Kragballe 2006 compared 3 52-week regimens with all treatments used once daily.

Combination therapy with calcipotriol and betamethasone dipropionate was associated with significantly lower rates of adverse events than either of the two 'complex' long-term regimens:

• alternating treatment with combination therapy for 4 weeks, then calcipotriol for 4 weeks (RD -0.08; 95% CI -0.17 to -0.00); and

• combination therapy for 4 weeks, then calcipotriol for 48 weeks (RD -0.16; 95% CI -0.25 to -0.08).

The difference between the two 'complex' regimens was not statistically significant at the 5% level (RD -0.08; 95% CI -0.17 to 0.01), although there was a trend in favour of alternating therapy.

Analysis 15: vitamin D analogues versus other treatment

A vitamin D analogue was compared with 12 other treatments (see Analysis 15.9 and Analysis 15.10). Data on local adverse events were available for 10 intervention-comparator contrasts, and systemic adverse events data were available for 8 intervention-comparator contrasts.

We assessed results from the analyses of adverse events data using a random-effects risk difference metric and report these separately. For local adverse events, three intervention-comparator contrasts were significantly different. These were the comparison of calcipotriol against calcipotriol + nicotinamide (RD -0.17; 95% CI -0.30 to -0.03), calcipotriol versus corticosteroid + salicylic acid (RD 0.09; 95% CI 0.02 to 0.15), and calcipotriol versus tacrolimus ointment (RD -0.19; 95% CI -0.37 to -0.01). No trial that reported the rate of systemic adverse events found a significant difference in this outcome.

Calcipotriol versus coal tar

Our review found no statistically significant difference for local or systemic adverse events.

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Calcipotriol versus coal tar polytherapy

Our review found no statistically significant difference for local or systemic adverse events.

Calcipotriol versus nicotinamide 1.4%, twice daily

Systemic adverse events data were not reported for this analysis. There was no significant difference between the local adverse event rates, although there was a trend in favour of calcipotriol (RD - 0.15; 95% CI -0.32 to 0.03; I² statistic = NA).

Calcipotriol versus calcipotriol + nicotinamide (0.05%, 0.1%, 0.7%, or 1.4%), twice daily

Systemic adverse events data were not reported for this analysis. Monotherapy with calcipotriol was significantly less likely to cause local adverse events than combined therapy with nicotinamide (RD -0.17; 95% CI -0.30 to -0.03; I² statistic = NA).

Calcipotriol versus corticosteroid + salicylic acid

Relative to calcipotriol alone, combination therapy with betamethasone dipropionate and salicylic acid was significantly less likely to be associated with local adverse events (RD 0.09; 95% CI 0.02 to 0.15). There was no significant difference in the rate of systemic adverse events.

Calcipotriol versus propylthiouracil cream

Our review found no statistically significant difference for local or systemic adverse events.

Calcipotriol versus tacrolimus ointment

Relative to calcipotriol, tacrolimus was associated with a significantly higher rate of local adverse events (RD -0.19; 95% CI -0.37 to -0.01). There was no significant difference in the rate of systemic adverse events.

Calcipotriol versus tazarotene

Our review found no statistically significant difference for local or systemic adverse events.

Calcipotriol versus tazarotene gel plus mometasone furoate cream

Adverse events data were not reported for this analysis.

Calcipotriol versus vitamin B12 cream

Our review found no statistically significant difference in the rates of local adverse events. Data on systemic adverse events were not reported.

Head-to-head vitamin D alone or in combination: dosing

Our review found no statistically significant difference for local or systemic adverse events.

Head-to-head vitamin D alone or in combination: occlusion

Local adverse events data were not reported for this analysis. Our review found no statistically significant difference in the rate of systemic adverse events.

Analysis 16: Flexural/facial psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of four topical treatments for inverse or facial psoriasis. Data on local adverse events were available for all four treatments, but data on systemic events were reported only for one treatment, i.e. the comparison of tacrolimus against placebo (see Analysis 16.9 and Analysis 16.10). Relative to placebo, participants receiving topical tacrolimus were significantly less likely to report local adverse events (RD -0.17; 95% CI -0.30 to -0.03), but there was no significant difference in the rate of systemic events. No other treatment was significantly different from placebo.

Analysis 17: Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for flexural or facial psoriasis, where vitamin D was compared with an active control. We identified five intervention-comparator contrasts (see Analysis 17.9 and Analysis 17.10). Local adverse events data were missing for the comparison of calcitriol and tacrolimus. There were no data on systemic events for any of the five contrasts. In people treated with calcipotriol, the rate of local adverse events was significantly higher compared to those receiving combination therapy with calcipotriol and hydrocortisone for facial psoriasis (RD 0.15; 95% CI 0.08 to 0.23) and also compared to those receiving calcitriol (RD 0.09; 95% CI 0.02 to 0.17). These findings aligned with the withdrawal rates for adverse events (Analysis

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17.7). We found no other significant differences in adverse events rates.

Analysis 18: Scalp psoriasis: placebo-controlled trials

This comparison included placebo-controlled trials of 11 topical treatments for scalp psoriasis (see Analysis 18.9 and Analysis 18.10). There were no adverse events data for two treatments, betamethasone valerate and amcinonide. Data on systemic events were available for four treatments.

Relative to placebo, participants treated with betamethasone dipropionate were significantly less likely to suffer local adverse events (RD -0.07; 95% CI -0.13 to -0.01; I² statistic = 0%). These findings aligned with the withdrawal rates for adverse events (Analysis 18.7). We found no other significant differences in adverse events rates.

Analysis 19: Scalp psoriasis: vitamin D alone or in combination versus other treatment

This comparison included head-to-head trials of treatments for scalp psoriasis in which one of the interventions was a vitamin D product (used either as monotherapy or in combination with another product) (see Analysis 19.9 and Analysis 19.10). We identified six intervention-comparator contrasts, and there were no adverse events data.

Calcipotriol monotherapy was associated with a significantly higher rate of local adverse events than five of the comparator treatments; these included betamethasone dipropionate (RD 0.07; 95% CI 0.04 to 0.11; I² statistic = 0%), betamethasone valerate (RD 0.17; 95% CI 0.01 to 0.33; I2 statistic = 76.5%), and clobetasol propionate (RD 0.19; 95% CI 0.10 to 0.28; I² statistic = 0%). Relative to calcipotriol, combined therapy with calcipotriol and betamethasone dipropionate (RD 0.09; 95% CI 0.06 to 0.12; I² statistic = 28.1%), and coal tar polytherapy (RD 0.24; 95% CI 0.15 to 0.33; I² statistic = NA), were also associated with significantly lower rates of adverse events. Compared with betamethasone dipropionate monotherapy, the rate of adverse events for combined therapy with calcipotriol and betamethasone dipropionate was not significantly different (RD -0.00; 95% CI -0.02 to 0.01; I² statistic = NA). There were no significant differences in the rates of systemic adverse events.

(ii) Findings from the separate search for additional studies of adverse events

In addition to findings on adverse events from the main review, we undertook a separate search for additional safety and tolerability studies. The update searches in 2011 identified 3 new literature reviews and 12 potentially relevant records. Eight studies (reported in seven references) met the inclusion criteria, and we excluded five studies from the review (Breneman 2007; Carboni 2005; Feldman 2007a; Hong 2010; and Hong 2011; Jacobi 2008) in addition to the 13 trials excluded in the previous edition of this review (Aste 2004; Bos 2002; Floden 1975; Franssen 1999; Kang 1998; Lebwohl 1996; Park 2002; Senter 1983; Singh 2000; Stevanovic 1977; Traulsen 2003; Uhoda 2003; Vissers 2004). Table 24 details the characteristics and key findings of the included studies, and Table 25 lists the excluded studies. Langner 1996 and van de Kerkhof 1996c also reported the study by Gerritsen 2001. Two papers reported two studies (Berth-Jones 1993; Kimball 2008), which we analysed separately. Kimball 2008 was a review reporting findings from phase II and phase III trials of clobetasol foam (we did not locate any other reports of these studies). The study by van de Kerkhof 2002c reported a two-phase open study of tacalcitol, where 'responders' to part 1 were eligible for part 2. Full results for part 2 were not reported separately (e.g. incidence of local adverse events was reported only for all trial participants). The study by Lambert 2002 appeared very similar to the second phase of the study reported by van de Kerkhof 2002c, but as there was no explicit evidence of an association, we analysed it as a separate trial.

Of the 29 included adverse events studies, 20 were uncontrolled (Barnes 2000; Berth-Jones 1992c; Berth-Jones 1993; Bleiker 1998; Brodell 2011b; Cullen 1996; Gerritsen 2001 and Langner 1996; Kimball 2008 (phase II trial); Kragballe 1991b; Lambert 2002; Menter 2007; Miyachi 2002; Poyner 1993; Ramsay 1994; Roelofzen 2010; van de Kerkhof 1997b; van de Kerkhof 2002c; Vazquez-Lopez 2004; Veraldi 2006; Wishart 1994), 8 were randomised trials (Andres 2006; Corbett 1976; Guzzo 1996; Katz 1987b; Katz 1989; Kimball 2008 (phase III trial); Lebwohl 1998b; Lebwohl 2001), 1 was a retrospective controlled study (Heng 1990), and 1 reported a control group of people using treatment other than calcipotriol (Berth-Jones 1993).

We did not include any of the eight RCTs in the main review. Six RCTs did not meet the inclusion criteria for the main review, because they did not address comparisons of interest (Andres 2006; Corbett 1976; Katz 1987b; Katz 1989; Lebwohl 1998b) or effectiveness (Guzzo 1996). We excluded one RCT from the main review because it did not report the numbers of participants in each arm of the trial (Lebwohl 2001). In all eight RCTs, the adequacy of concealment of treatment allocation was unclear, the method of randomisation was unclear, and all were double-blind (participant/investigator) trials. Five of the eight trials reported baseline comparability, with two studies demonstrating betweengroup comparability in both clinical and demographic characteristics (Katz 1987b; Katz 1989), two reporting differences in baseline severity (Guzzo 1996; Lebwohl 1998b), and one with some significant differences in demographics and clinical severity (Andres 2006). Four of the eight RCTs were placebo-controlled and six included active controls.

Of the nine controlled studies, two were within-patient designs (Corbett 1976; Katz 1989), and the remainder were betweenpatient (parallel-group) designs.

Trials ranged in size from 10 to over 4000 participants. Treatment

duration ranged from 2 weeks to 18 months. Loss to follow up averaged 9.5% (range = 0% to 63%). The mean age of participants was 48 (range = 15 to 85), and participants were more likely to be men (mean: 55%; range = 40 to 82%). In 12 studies, the baseline severity of participants was unclear. Remaining studies recruited participants with mild to moderate disease (N = 4), mild to severe disease (N = 4), moderate to severe disease (N = 6), and severe disease (N = 3).

Study treatments included vitamin D products (18/29), topical corticosteroids (12/29), coal tar (1/29), and tazarotene (2/29). Some comprised of combination regimens, such as vitamin D and corticosteroid (Lebwohl 1996; Lebwohl 1998b; Vazquez-Lopez 2004) or tazarotene and corticosteroid (Lebwohl 2001). No study of dithranol met the inclusion criteria for the review. Twenty-one of the studies assessed local adverse events, 18 assessed systemic effects, and 11 studies assessed both types. Although six RCTs reported some data on cutaneous adverse events (Andres 2006; Corbett 1976; Katz 1987b; Katz 1989; Lebwohl 1998b), these were neither suitable nor adequate for pooling.

Vitamin D products (N = 18)

Eleven studies evaluated local (N = 7) or systemic (N = 8) adverse effects associated with calcipotriol, or both. The rate of withdrawals due to local adverse events ranged from 4% to 14% and the rate of adverse events ranged between 20% and 41%; larger trials reported higher rates (weighted mean: 36%). In the 52-week trial by Barnes 2000, facial irritation affected 30% of participants in the early stages of the trial, but the incidence declined over time. The incidence of systemic effects was less common: 5/8 studies found no significant effects. Bleiker 1998's study of inpatients with severe disease found 5/28 developed hypercalcaemia after receiving a dose greater than 5 g/kg. The study by Berth-Jones 1993, in which 10 trial participants received a weekly dose of 100 g of calcipotriol for 4 weeks, found that urinary calcium increased significantly from baseline levels.

Four studies evaluated both local and systemic adverse effects associated with tacalcitol. The rate of withdrawals due to local adverse events ranged from 0% to 6%, and the rate of adverse events ranged between 10% and 21%. Three studies found no systemic effects. The study by van de Kerkhof 2002c found that over half of study participants with psoriasis affecting 10% to 20% of their body surface area exceeded the recommended daily dose of 5 g/day (up to 13 g daily), but there was no effect on calcium homeostasis. Systemic effects were identified in over half enrolled participants in the trial by Miyachi 2002, but only 6/155 events were considered to be treatment-related in this uncontrolled study.

Three studies looked at adverse events associated with calcitriol (Brodell 2011b; Gerritsen 2001; Wishart 1994). One study examined the tolerability and systemic effects of calcitriol used as monotherapy (3 mcg/g ointment applied twice daily, as per licence) (Gerritsen 2001). Three per cent of participants withdrew due to adverse events, and 15% reported local adverse events. The withdrawal rate due to systemic effects was low (0.4%), but 4 cases of hypercalcaemia were reported (N = 253). Mean daily use of calcitriol in this trial was 6 g (range = 1 to 24 g). In Brodell 2011b, participants who had responded to treatment with clobetasol spray then received 8 weeks of maintenance treatment with calcitriol 3 mcg/g ointment twice-daily. Around 15% (35/235) of participants reported burning or stinging at the end of treatment. Wishart 1994 tested the effects of high-dose calcitriol (15 mcg/ g once-daily) on 3 groups of participants, with the quantity used proportional to the area affected The trial recruited participants with psoriasis affecting up to 30% of the body surface area. Mean daily drug use ranged from 74 to 306 mcg. The study did not observe any systemic adverse events, skin irritation, or 'clinically relevant' changes in vital signs, haematology, biochemistry, urine, or electrocardiograms.

Corticosteroids (N = 12)

The 12 studies of adverse events associated with steroids adopted a range of different study designs. Four studies had no comparator group (Brodell 2011b; Kimball 2008 (Phase II); Menter 2007; Vazquez-Lopez 2004). Seven studies were randomised trials (Andres 2006; Corbett 1976; Katz 1987b; Katz 1989; Kimball 2008 (Phase III); Lebwohl 1998b; Lebwohl 2001), of which six used active treatment controls and three were placebo-controlled. The remaining study retrospectively compared 13 participants who had used topical corticosteroids for between 6 months and 12 years with a 'no steroid' group (N = 15).

Eight of the 12 corticosteroid studies assessed atrophy or preatrophy in people with psoriasis who were treated with topical steroids. Three studies explicitly described the methods used to assess atrophy, and some studies did not clearly state the numbers of participants affected by atrophy or its extent.

The retrospective study by Heng 1990 compared 13 participants who had used topical corticosteroids for between 6 months and 12 years with a 'no steroid' group: These 15 individuals had previously used tar, UVB, or were untreated. Light microscopy revealed no between-group differences. However, electron microscopy revealed multi-layered, fragmented, and disorganised basal laminae (the lining of the outer surface of the cell membrane) in the steroid group; this appeared to be correlated with duration of treatment. Fragmentation was not observed in the control group.

The 4-week study by Katz 1989 identified preatrophy in 20% of involved plaques, using a hand-held magnifying lens. Two longterm (26-week) studies of combination maintenance therapy with steroids (Lebwohl 1998b; Lebwohl 2001) did not observe cutaneous atrophy, but neither study reported the assessment method. A 4-week trial of clobetasol propionate for scalp psoriasis assessed atrophy using an ultrasound probe (20 MHz) (Andres 2006). No case of cutaneous atrophy was detected in either the gel (which was left in) or shampoo formulations (which was rinsed out after 15 minutes). However, there was a significantly greater reduction in skin thickness in participants treated with the gel, compared to the shampoo group (Andres 2006).

An uncontrolled study of clobetasol spray (Brodell 2011b) detected atrophy in 7/285 participants after 4 weeks of treatment, though the study did not describe the assessment method. A randomised trial compared 2 weeks of treatment with clobetasol propionate foam (N = 253), clobetasol propionate ointment (N = 121), or placebo foam (N = 123) (Kimball 2008, Phase III). The paper did not describe the assessment method. The trial found five cases of skin atrophy in the clobetasol foam group and one case in the placebo group, but the paper did not report the number of atrophy cases in the clobetasol ointment group. COBRA (Clobex Spray Community-Based Research Assessment) was an open-label 4-week study where 1421 people with psoriasis used clobetasol spray 0.05% twice daily as monotherapy (Menter 2007). Cases of telangiectasia, skin atrophy, or folliculitis each occurred in less than 1% of participants (the study did not report the numbers affected or the assessment method).

Four studies examined systemic effects associated with corticosteroids. The study by Corbett 1976 compared betamethasone valerate with clobetasol propionate. Quantities used by study participants were small (mean: 7 g/wk), and the study observed no pituitary-adrenal suppression. Katz 1987b compared two 'superpotent' corticosteroids and identified temporary and reversible adrenal suppression in 20% (8/40) of study participants. In the study by Andres 2006, none of the 14 participants assigned to clobetasol propionate shampoo experienced hypothalamic-pituitaryadrenal (HPA) axis suppression. Conversely, 2 of the 12 participants in the gel group experienced HPA axis suppression. Neither formulation had an impact on ocular safety. In a phase II study (Kimball 2008), higher levels of clobetasol were found in the plasma tests in ointment group than in the shampoo group, though the difference was not statistically significant.

Tazarotene (N = 2)

The study by Lebwohl 2001 compared three types of maintenance therapy: tazarotene plus steroid, tazarotene plus placebo, and placebo alone. In this 6-month trial, there were no withdrawals due to adverse events, but 24% of participants in the tazarotene/ steroid group and 29% of participants in the tazarotene/placebo group experienced adverse events. There were no adverse events in the placebo group, and the study did not assess systemic effects. Veraldi 2006 evaluated short-contact treatment with tazarotene gel in 43 participants. The study drug was applied once daily, left for 20 minutes, and then rinsed off with water. The number of participants reporting pruritis and burning decreased over the 45day study period, with 14/43 participants reporting mild pruritis at the end of treatment, and the same number reporting mild burning. The study did not assess systemic effects.

(c) Quality of life measures

Eight of the 177 studies included in the main review formally assessed quality of life (QoL) (Alora-Palli 2010; Bernstein 2006; Cook-Bolden 2010; Guenther 2002 (H) and Guenther 2002 (P); Hutchinson 2000; Saraceno 2007; Van de Kerkhof 2006; Wall 1998).

The trials used different QoL measures. Alora-Palli 2010 used the Dermatology Quality of Life Index (DLQI), and Bernstein 2006 used the Quality of Life Index. In both measures, higher scores indicate poorer QoL. The trial by Guenther (2002) measured quality of life using the Psoriasis Disability Index (PDI) and the Euro-QOL (EQ-5D and EQ-VAS), reporting it in a separate publication (van de Kerkhof 2004). Hutchinson 2000 and Wall 1998 also assessed quality of life using the Psoriasis Disability Index; Wall 1998 also used the Sickness Impact Profile (SIP). Although four studies used the PDI to measure quality of life, they did not report data in sufficient detail to allow pooling, so we reported findings narratively. Four trials included participants with mild to moderate disease; the other three trials included participants with at least moderately severe disease. The number of participants ranged from 60 (Alora-Palli 2010) to 828 (Guenther 2002 (P)), although PDI scores were obtained for only 51% of participants in this trial (van de Kerkhof 2004). Saraceno 2007 and Van de Kerkhof 2006 used the Skindex-29, with Van de Kerkhof 2006 also employing the SF-36. One trial used a QoL instrument designed specifically for the scalp, the Scalpdex (Cook-Bolden 2010).

Dermatology Quality of Life Index (DLQI) (N = 1)

The Dermatology Quality of Life Index (DLQI) is scored from 0 to 30, where higher scores indicate poorer QoL. Alora-Palli 2010 used the DLQI to compare calcipotriol and liquid carbonis distillate (LCD) 15% solution. Both treatments improved QoL relative to baseline, but there was no significant difference between the groups at the end of the 12-week treatment period. Participants were followed up for 6 weeks post-treatment; in the LCD group, quality of life continued to improve, but it deteriorated in the calcipotriol group. There was a significant between-group difference in the DLQI scores at the 18-week follow-up assessment. This QoL effect reflected the PASI and physician global assessments, which demonstrated significantly lower recurrence rates in the LCD group at 18 weeks.

EuroQOL (EQ-5D and EQ-VAS) (N = 1)

The EQ5D is a generic quality of life measure. van de Kerkhof 2004 reported quality of life data from the trial by Guenther 2002. The study presented the EQ5D scores in a non-standard method, making their interpretation problematic. EQ-VAS (scored from 1 to 100) supported findings from the PDI assessments, with all groups improving quality of life scores relative to baseline. However, the EQ-VAS score for the once-daily combined calcipotriol/betamethasone dipropionate (plus placebo) group was

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higher (with a better quality of life) than the corresponding score in the twice-daily group. Relative to baseline, improvements in mean quality of life scores were statistically significant in all treatment groups, but the significance of between-group differences was not reported.

Psoriasis Disability Index (PDI) (N = 4)

In all four trials reporting this measure, there was an improvement in mean quality of life scores for participants in every group. The trial by Guenther 2002 found the greatest improvement (reduction in PDI from baseline, i.e. improvement in QoL) for twicedaily combination treatment with calcipotriol and betamethasone (50%). Corresponding figures were as follows: once-daily combination treatment with calcipotriol and betamethasone (plus placebo): 41%, calcipotriol twice daily: 31%, and placebo twice daily: 9%. In terms of absolute scores, the group using twice-daily combination treatment with calcipotriol and betamethasone had the best (lowest) quality of life score, followed by the once-daily combined treatment group, calcipotriol group, and placebo group. Relative to baseline, improvements in mean quality of life scores were statistically significant in all treatment groups, but the statistical significance of between-group differences was not reported. In the study by Hutchinson 2000, quality of life improved from baseline significantly more in the group treated with calcitriol relative to the dithranol group. The comparison of dithranol and calcipotriol by Wall 1998 found that the magnitude of the difference was greater in the calcipotriol group, but the difference was not statistically significant at the 5% level.

Quality of Life Index (N = 1)

Bernstein 2006 used the Quality of Life Index to compare *Mahonia aquifolium* with placebo in participants with mild to moderate psoriasis. The index ranges from 0 to 120, with higher scores indicating poorer quality of life. *Mahonia aquifolium* was significantly more effective than placebo, and this was also reflected in the QoL index.

Scalpdex (N = 1)

Cook-Bolden 2010 used the Scalpdex, a scalp dermatitis-specific quality of life instrument comprising 23 questions, each scored 0 (never) to 100 (all the time). Relative to baseline scores, the study found a significant improvement in QoL for participants in the clobetasol spray arm of the trial and no significant improvement in the placebo spray arm.

SF-36 (N = 1)

The SF-36 is a generic short-form health survey with 36 questions covering physical and mental health. The study by Van de Kerkhof 2006 compared calcipotriol and dithranol administered in a day-

care setting. The study found no significant difference in quality of life, either for the Skindex-29 or for the SF-36.

Sickness Impact Profile (SIP) (N = 1)

The SIP is a generic quality of life measure. Wall 1998 assessed participants using the SIP, which has a maximum score of 136. As with the PDI, the SIP found a statistically significant improvement from baseline in both groups, but the between-group difference was not statistically significant.

Skindex-29 (N = 2)

The Skindex-29 is a dermatology-specific QoL index that includes three domains: emotions, symptoms, and functioning. Two trials used the Skindex-29, both enrolling participants with mild to moderate disease. Saraceno 2007 compared a 12-week course of calcipotriol against 4 weeks of combination therapy (calcipotriol/ betamethasone dipropionate) followed by maintenance with calcipotriol for 8 weeks. Quality of life improved in both groups relative to baseline. Van de Kerkhof 2006's study in a day-care treatment setting found no significant difference between calcipotriol and dithranol, either for the Skindex-29 or for the SF-36.

(d) Economic outcomes

(THIS SECTION HAS NOT BEEN UPDATED)

A number of studies have looked at economic aspects of topical treatment for psoriasis. These include cost-of-illness studies (Feldman 1997; Jenner 2002; Poyner 1999), quality-of-life studies (Leu 1985; Lundberg 1999; Ortonne 2000; Schiffner 2003; Zachariae 2002; Zug 1995), methodological issues (Lambert 1996; Lambert 1999), willingness to pay analyses (Lundberg 1999; Poyner 2000; Schiffner 2003), cost analysis (Feldman 2000), and cost-effectiveness analyses (Ashcroft 2000; de Tiedra 1997; Harrington 1995; Köse 1997; Marchetti 1998; Oh 1997; Owen 1993; Parodi 1991; Schwicker 1992; Sorensen 2002; Stern 1988). These analyses involve a range of modelling approaches and assumptions and have not been formally reviewed here, although they were part of our original review (Mason 2002a). Our updated review revealed no substantial variations in tolerability or effectiveness for most treatment comparisons, and no trials provided robust resource data on the consequences of managing treatment failure. Consequently, any economic models extrapolating beyond the duration of the trials may be largely speculative and uninformative. In the light of available data, a 'cost and consequences' approach, in which costs and outcomes are reported separately, may be most informative to clinical decision-makers. The relative short-term clinical performance of topical antipsoriatic treatments can be set against their reimbursed costs. While it is accepted that long-term sequelae in participants not responding to treatment

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may be very important when considering overall costs and benefits, there are no good comparative data on these costs with which to distinguish between treatments (Mason 2002a).

(e) Concordance or adherence with treatment

(THIS SECTION HAS NOT BEEN UPDATED)

The separate search for studies of concordance/adherence identified 246 potentially relevant studies for screening. Of these, we retrieved 18 papers, and we included in the review 12 papers reporting on 11 studies (see Table 26) (Balkrishnan 2003; Carroll 2004a; Carroll 2004b; Feldman 2007; Ferrandiz 1998; Fouere 2005; Gokdemir 2008; Richards 1999; van de Kerkhof 1998c; van de Kerkhof 2000; van de Kerkhof 2001; Zaghloul 2004). Some studies were pilots (e.g. Balkrishnan 2003; van de Kerkhof 1998c) for other studies (i.e. Carroll 2004a; van de Kerkhof 2000). We listed studies that did not meet the inclusion criteria in Table 27 (Atkinson 2004; Chu 2000; Gupta 2007; Lee 2006; Osborne 2002; Richards 2006; Szeimies 2004). We identified three reviews and bibliographies were checked for further potentially relevant studies (Gupta 2007; Lee 2006; Richards 2006). Of the 11 studies included in the review, 2 were randomised controlled trials (Carroll 2004a; Ferrandiz 1998) and 4 were questionnaire surveys Fouere 2005; Richards 1999; van de Kerkhof 1998; van de Kerkhof 2000). The remaining five studies were prospective experiments, of which one included a control group (van de Kerkhof 2001). The number of participants included in the adherence studies ranged from 10 to 1281, and the study duration ranged from 1 to 16 weeks. In total, 11 studies enrolled 5541 participants, of which 39% were men and the mean age was 47.0 (range = 4 to 91). Study size ranged from 10 to 1281 participants, with the questionnaire surveys reporting the largest sample sizes. Studies covered four main issues. First, some reported methodological issues on the measurement of adherence (Balkrishnan 2003; Carroll 2004a; Fouere 2005; Zaghloul 2004). Second, some studies reported adherence rates (Balkrishnan 2003; Carroll 2004a; Fouere 2005; Richards 1999; van de Kerkhof 2000; van de Kerkhof 2001; Zaghloul 2004). Third, seven studies considered reasons for non-adherence (Carroll 2004a; Gokdemir 2008; Fouere 2005; Richards 1999; van de Kerkhof 1998c; van de Kerkhof 2000; Zaghloul 2004). Fourth, three studies assessed interventions to improve adherence (Feldman 2007; Ferrandiz 1998; van de Kerkhof 2001).

Methodological issues relating to adherence measurement and adherence rates

Some studies used self-reported adherence rates (Gokdemir 2008; Richards 1999; van de Kerkhof 2000; van de Kerkhof 2001; Zaghloul 2004). In these studies, which included short-term prospective trials and cross-sectional questionnaire surveys, rates varied between 61% and 72%. However, other studies adopted more objective assessment methods. The study by Fouere used a semistructured participant questionnaire: The PMAQ-3w scale (Patient Medication Adherence Questionnaire) asked about strict adherence to prescribed regimen over the previous three days and previous weekend. Using this method, the study (Fouere 2005) deemed just 27% of participants as 'compliant'. Some studies adopted a single-blind approach to assess electronic bottle caps (i.e. participants were unaware that bottles were fitted with electronic measuring devices) (Balkrishnan 2003; Carroll 2004a; Feldman 2007). Known as a 'MEMS cap' (Medication Event Monitoring System), the medication bottle cap was fitted with a microprocessor to record the time/date of every opening of the bottle. When compared with participant logs, electronic methods suggested that adherence was considerably lower than rates reported by participants. For example, Balkrishnan's 1-week pilot study of 10 participants found that the adherence rate was 67% by electronic assessment compared with a self-reported rate of 92%. Carroll and colleagues took this pilot study forward using an 8-week trial of 30 participants (Carroll 2004a). Adherence rates fell over time and averaged 55% over the study period when assessed using the MEMS cap. Twice-daily dosing was achieved on 39% of the treatment days.

Reasons for non-adherence

Reasons for non-adherence comprised therapy characteristics (real or perceived), participants' clinical characteristics, and participants' demographic characteristics. Regarding therapy characteristics, the included studies identified the following as influences on adherence: efficacy (Fouere 2005; van de Kerkhof 1998c; van de Kerkhof 2000), side-effects (Fouere 2005; Richards 1999; Zaghloul 2004), time for application (Fouere 2005; Gokdemir 2008; van de Kerkhof 2000), and 'messiness' (Fouere 2005; Richards 1999; van de Kerkhof 1998c). Adherence was higher for topical therapy compared with systemic treatment (Zaghloul 2004), but adherence rates also varied by type of topical treatment (Fouere 2005). Disease severity was inversely related to adherence (Carroll 2004a; Richards 1999), although the direction of causality is unclear. Higher adherence rates were variously associated with female gender (Carroll 2004a; Zaghloul 2004), older age (Carroll 2004a; Richards 1999), higher educational achievement (Gokdemir 2008), and older age of disease onset (Richards 1999), but findings on marital status were mixed (Gokdemir 2008; Zaghloul 2004).

Methods to improve adherence

Office visits to clinicians were found to temporarily increase adherence rates (Carroll 2004a; Feldman 2007). An educational programme designed to improve adherence had no impact on the effectiveness of treatment (Ferrandiz 1998). Van de Kerkhof's study comparing ointment/cream regimen with ointment-only treatment found no clear impact, with 51% of participants reporting better adherence with the cream/ointment option (van de Kerkhof 2001).

DISCUSSION

Summary of main results

Effectiveness

This review analysed evidence from 177 studies that included 34,808 participants. We reported results separately for treatments of the body, inverse psoriasis, and psoriasis of the scalp.

Treatments for psoriasis of the body

We analysed six placebo-controlled comparisons of treatments for psoriasis of the body. Most treatments were significantly more effective than placebo, with the equivalent pooled effect on a 6point improvement (IAGI) scale ranging from 1.02 (potent corticosteroid) to 1.80 (very potent corticosteroid). Pooled effects indicated similar improvements for vitamin D analogues (1.03 points), dithranol (1.22 points), and combination therapy with vitamin D and potent corticosteroid (1.65) on a 6-point scale.

Amongst less frequently researched treatments, 13 were significantly more effective than placebo. For these treatments, the equivalent effect on a 6-point IAGI scale ranged from 0.55 (calcipotriol plus nicotinamide) to 3.40 (herbal skin care products). Tazarotene achieved an equivalent improvement of 0.99 on the 6-point IAGI scale. One small within-patient study found that tar was no more effective than placebo (Kanzler 1993). Findings from single studies should be treated with caution and regarded as requiring further evidence before guiding treatment.

We evaluated nine head-to-head comparisons. Findings from the comparisons of vitamin D against potent or very potent corticosteroids were mixed, and they varied depending on which vitamin D and which corticosteroid were analysed. Combination treatment with vitamin D/corticosteroid was usually more effective than monotherapy with the same vitamin D, and it was always more effective than the same corticosteroid used alone.

Twelve complex regimens (i.e. treatment sequences that do not consist of a simple head-to-head comparison between two active treatments) were compared with vitamin D. Seven of the complex regimens were more effective than vitamin D, achieving an additional benefit of between 0.27 points and 0.72 points on an equivalent 6-point IAGI. A comparison of three long-term regimens found no significant difference in effect, but some were better tolerated (see below). These studies were interpreted independently, reflecting the differing management regimens. None of the studies presented have been replicated precisely. Therefore, while supportive evidence for findings can be drawn from similar studies, trial findings for complex regimens should be interpreted with caution when informing clinical decisions.

Treatments for inverse psoriasis

The review included four placebo-controlled comparisons of treatments for psoriasis of the sensitive areas of the body. As there were no effectiveness data for tacrolimus, the review evaluated benefits for three treatments. All three were significantly more effective than placebo, with the pooled effect on a 6-point IAGI scale ranging from 0.98 (pimecrolimus) to 3.25 (betamethasone valerate). Five treatments were compared head-to-head against vitamin D. Betamethasone valerate, combined treatment with calcipotriol and hydrocortisone, and calcitriol were all significantly more effective than calcipotriol alone. On an equivalent 6-point IAGI, these treatments achieved an additional benefit of 2.22, 0.33, and 0.67 points, respectively.

Treatments for psoriasis of the scalp

We compared 11 treatments for scalp psoriasis with placebo. All but two were significantly more effective than vehicle alone; the most effective was the very potent corticosteroid, clobetasol propionate, which delivered a benefit of almost 2 points (1.88) on an equivalent 6-point IAGI scale. When compared head-to-head, vitamin D was significantly less effective than both potent and very potent corticosteroids. On an equivalent 6-point IAGI scale, the additional benefit was between 0.48 and 0.67 points. Combination therapy with calcipotriol and betamethasone dipropionate was significantly more effective than either product used alone.

Adverse effects

Randomised evidence found that vitamin D was significantly more likely than potent corticosteroids to cause local adverse effects (15% versus 8%, P = 0.006), and participants were (borderline) more likely to withdraw for this reason (2% versus 1%, P = 0.059). These findings were supported by indirect comparisons of placebocontrolled trials. Participants tolerated combined treatment with vitamin D/corticosteroid on either the body or the scalp as well as potent corticosteroids and significantly better than vitamin D alone. No comparison of topical agents found a significant difference in systemic adverse effects.

Studies of longer-term adverse events found that up to 40% of those using vitamin D analogues experienced cutaneous effects and that up to 14% of participants stopped using vitamin D analogues for this reason. There was some evidence to suggest that using vitamin D analogues at high doses could cause systemic adverse effects.

Two of the major cutaneous adverse effects of topical corticosteroids are dermal and epidermal atrophy (Hengge 2006; Kragballe 2006). Only 25 of the 78 RCTs of corticosteroids included in the review explicitly assessed cutaneous atrophy, and in these the duration of the trials, skin sites assessed, and methods employed reduced the chance of detection. The studies either did not support the methods used to assess atrophy (14/25 trials), physicians undertook them (10/25), study participants self-reported them (5/ 25), or a combination of the aforementioned. Where physician assessments were conducted, the methods used were not explicit (for example, whether microscopy or ultrasound was used). Two of the 25 trials featured an independent 'adjudication committee', which assessed whether atrophy was likely to be treatment-related (Kragballe 2006; Poulin 2010). Trial duration of the 25 trials was typically between 4 and 8 weeks, but there were 2 6-month studies (Katz 1991a; Poulin 2010) and 2 52-week studies (Kragballe 2006; Luger 2008).

Two trials that assessed atrophy did not report whether or not any cases of atrophy occurred (Huang 2009; Koo 2006). Ten RCTs reported cases of atrophy, and these were trials of either very potent corticosteroids (Cook-Bolden 2010; Decroix 2004; Lowe 2005; Poulin 2010; Reygagne 2005) or combination treatment with potent corticosteroids (Guenther 2002 (H); Kragballe 2004; Kragballe 2006; Papp 2003 (H) and Papp 2003 (P); Yang 2009). Three of the 10 were trials of scalp psoriasis (Cook-Bolden 2010; Poulin 2010; Reygagne 2005), all of which investigated clobetasol propionate.

Some studies reviewed as part of our search for additional studies of adverse events of longer-term use of topical corticosteroids revealed atrophy and damage to basal laminae, the extent of which appeared to be correlated with dosage and duration of use. Research of very potent steroids on the skin of normal volunteers (Kao 2003). Longer-term use with topical retinoids caused cutaneous effects in around 30% of participants, but systemic effects were not assessed.

Concordance/adherence

Thirty-four of the 177 RCTs assessed adherence ('compliance'), and 26 trials reported findings. Trials used various methods such as self-report by participants, medication weight, and treatment completion. Most of the 26 studies that reported findings on adherence found it to be suboptimal. However, Gupta has argued that the relevant notion is concordance rather than compliance or adherence (Gupta 2007). Whereas adherence implies that people comply with clinicians' prescribed treatment, concordance involves a negotiated doctor-patient agreement about the treatment regimen. Van de Kerkhof's survey of over 800 European people with psoriasis found that some chose not to comply, preferring to apply the 'minimum' dose needed to achieve the effect they wanted (van de Kerkhof 2000). If prescribed regimens reflected the wishes of the person with psoriasis, 'adherence' rates might be expected to rise. Nonetheless, poor adherence to treatment regimens is likely to impair the effectiveness of treatment.

The review identified one study that used an educational programme to improve adherence rates in people with psoriasis (Ferrandiz 1998). This study found no statistically significant difference in outcomes, a finding which contrasts with evidence concerning an educational programme for atopic dermatitis (Cork 2003). However, whereas the programme in the Ferrandiz 1998 study provided education about the disease, the study by Cork involved specialist nurses teaching methods for applying topical treatments. Therefore, the findings from Ferrandiz 1998 may not generalise to reflect the value of other educational programmes.

Sensitivity analysis

In four comparisons, we used sensitivity analysis to explore whether within-patient versus between-patient study designs were associated with different effect sizes. When comparing within-patient and between-patient study designs, our review found no statistically significant difference in treatment effect size. If withinpatient studies featured some correlation, due to systemic effects or cross contamination, then one would expect systematically smaller effect sizes from within-patient studies. If there is no correlation, then one would expect the variance to be smaller (as other sources of between-patient variation have been removed) but the effect size to be similar. We found no evidence to support a correlation for within-patient studies, and variances were similar for within and between-patient studies (Table 3), supporting the pooling of these, although it is accepted that within-patient designs are 'underweighted' in meta-analyses.

In the placebo-controlled studies of vitamin D products, there was little difference when comparing within-patient and between-patient study findings. On average, within-patient studies identified a slightly greater effect than between-patient studies (mean difference: 0.36 points on a 6-point IAGI scale). For placebo-controlled potent corticosteroids, there was a more pronounced difference in favour of within-patient studies (mean difference: 0.55 points on a 6-point IAGI scale). In the case of very potent corticosteroids, the mean effect relative to placebo was slightly larger in the betweenpatient studies (mean difference: 0.07 points on a 6-point IAGI scale), and the analysis of vitamin D against potent corticosteroid was similar in this respect (a small positive difference in favour of between-patient studies). These non-statistically significant differences may be chance findings.

Quality of the evidence

All included trials were randomised, but only 47 studies (27%) clearly reported the method used to randomise participants (Figure 3). Just 15 trials (9%) adequately concealed treatment allocation, but most (74%) masked participants to treatment. Of the 177 studies assessed, 151 reported loss to follow up, and 143 demonstrated that groups were comparable at baseline. Based on the 100 studies that reported assessable data on baseline severity, it was apparent that a wide range of severity was included within and between trials. If the PASI is unreliable for mild-to-moderate disease

(PASI scores less than 10) (Brownell 2007), it is possible that some trials may have contributed unreliable PASI data to this review. Trials varied in their treatment duration and in their outcome measures (see Included studies section for details). One reason for the variation in trial treatment duration is that the time taken for an intervention to be effective differs between treatments. The analyses and forest plots do not provide information on treatment duration, but this is an important consideration for clinical decisions. Trials also used different outcome measures; for example, only 113 studies reported data on the IAGI or IGA, and 65 studies reported PASI data. In order to maximise the number of studies contributing data for a particular treatment comparison, we used a combined end point that incorporated different outcome measures. This approach is not ideal; although all outcome measures essentially assess redness, thickness, and scaling (and sometimes area of psoriatic involvement), they differ in their construct. Therefore, the composite end points should be seen as indicative rather than definitive.

Medical text books commonly document the risk of skin atrophy and tachyphylaxis (diminution of the effects of a drug with continual use) as problems associated with topical corticosteroids (Bos 2008). Randomised evidence reported in this review suggests that maintenance regimens using intermittent dosing are safe and effective, and no RCT included in the review detected a statistically significant difference in systemic effects. However, the RCTs did not clearly report the methods employed (e.g. whether the investigator used ultrasound) and may have reduced the chance of detection. We undertook a separate search for longer-term studies of adverse events, which identified 12 relevant studies of corticosteroids. Studies were heterogeneous in terms of design, assessment methods, comparators considered, and doses used, so we did not pool data (Higgins 2011). Eight studies assessed atrophy, and six found some evidence of basal damage and atrophy. We identified no studies reporting adequate data on tachyphylaxis, despite including this term in our searches.

Potential biases in the review process

There are a number of limitations to this review. First, one author extracted data and a second checked it. Ideally, two authors should extract data independently (NHSCRD 2001). However, this approach was not feasible for this review because of funding constraints. Second, requests for unpublished data from trialists and sponsors were of variable success; requests were more likely to be successful when made for more recently published studies, products still on patent, or both.

This review included 19 comparisons with 10 potential outcomes, so multiple analyses may be expected to generate occasional chance findings. Thus, greater importance is attached where supportive findings are found from placebo-controlled and head-to-head studies as well as linked therapeutic modalities.

Agreements and disagreements with other studies or reviews

A systematic review by Ashcroft and colleagues focused on headto-head trials of one vitamin D analogue, calcipotriol (Ashcroft 2000a). Based on data from 37 studies with over 6000 participants, the review found that calcipotriol was at least as effective as potent topical corticosteroids and more effective than calcitriol, tacalcitol, coal tar, and short contact dithranol. Our review generally supported these findings. However, we found no significant difference for the comparison of calcipotriol and calcitriol (SMD -0.41; 95% CI -1.46 to 0.64), a finding based on 2 head-to-head trials with 261 participants. Our review also found that calcitriol is more efficacious than calcipotriol when used for inverse psoriasis, and it is also better tolerated. Our findings on the effectiveness of short-contact dithranol compared with calcipotriol were mixed, but physicians and people with psoriasis may be interested to know that inpatient treatment with dithranol appears more effective than calcipotriol (Monastirli 2000). Our review also supported findings from the review by Ashcroft 2000a that although calcipotriol caused significantly more skin irritation than potent topical corticosteroids, skin irritation rarely led to withdrawal of calcipotriol treatment.

The review by Afifi 2005 concluded that combined treatment with steroids and vitamin D analogues or topical retinoids appeared the most promising current treatment on account of its superior efficacy and favourable side-effects profile. However, longer-term adverse effects were not addressed by this review. In contrast, the review by Bruner 2003a focused on adverse effects. This review supported our findings that there is little robust evidence on longer-term adverse effects; therefore, concluding that since clearance is not a realistic expectation, "reasonable goals" are needed to avoid increasing the risk of cutaneous and systemic side-effects through over use.

AUTHORS' CONCLUSIONS

Implications for practice

Evidence from large numbers of trials indicates that most of the topical treatments tested in the trials reviewed here alleviate the symptoms of psoriasis. However, it was not possible to assess the performance of treatments at different levels of severity of psoriasis.

The evidence suggests that vitamin D products are more effective than emollient alone. Potent and very potent corticosteroids are also effective, and very potent corticosteroids are more effective than either potent corticosteroids or vitamin D products. The effectiveness of dithranol and tazarotene appears to be similar to that of vitamin D products. Although vitamin D and corticosteroids are equally effective for treating psoriasis of the body, corticosteroids appear to be more effective than vitamin D for treating psoriasis of the scalp. Combined treatment of vitamin D with corticosteroid is more effective than either vitamin D alone or corticosteroid alone. Vitamin D is more effective than coal tar, but findings on the relative effectiveness of vitamin D and dithranol were mixed. Occlusion enhances the effectiveness of vitamin D, as does twice-daily rather than once-daily application.

Compared with vitamin D alone, combined therapy that uses two products separately (vitamin D in the morning and corticosteroid at night) can achieve similar effects and be as well tolerated as using a combined product. However, some corticosteroids seem to perform better than others when used separately (see Analysis 12.5 and Analysis 12.9), and disease severity may also affect treatment performance, though poor reporting of baseline severity in trials means that we cannot confirm this. Use of a combined product may also enhance concordance, and there is evidence to suggest that adherence is higher when application time is shorter.

Potent corticosteroids are less likely than vitamin D to cause local adverse events, and treatment with corticosteroids is less likely to result in discontinued use because of these adverse events. Tazarotene is more likely than emollient to cause local adverse events. Our review found no difference between placebo and any other topical treatment in the assessment of systemic adverse events. However, this may reflect an absence of evidence (trials failing to appropriately assess these events over adequate time periods) rather than being evidence of absence.

Although current evidence demonstrates that topical steroids are as effective as and at least as well tolerated as vitamin D analogues, concern remains about the potential safety problems associated with corticosteroids (Bos 2008). Concerns include the risk of rebound (a worsening of disease following treatment discontinuation), skin atrophy (skin thinning), and tachyphylaxis (decreasing response to the drug) after long-term use (Hengge 2006). Methods to assess rebound have been developed and should be used in future research (Carey 2006). Regarding skin thinning, one problem with psoriasis is that the skin is very thick and a goal of therapy is to reduce the thickness of lesional (epidermal) skin. Damage to the surrounding normal skin may occur, and for that reason, people should use topical corticosteroids for limited periods or sparingly in delicate areas, such as the face or folds of the skin.

Although topical corticosteroids have been in use for about 50 years, there is a surprising lack of relevant evidence addressing steroid-associated skin dermal atrophy in people with chronic plaque psoriasis requiring long-term treatment. It is improbable that short-term (less than three weeks) courses of topical corticosteroids cause dermal skin atrophy, except in delicate areas such as the face and flexures (groin, axillae, inframammary). However, treatment may be long-term and continuous for people with more severe chronic psoriasis. Current assessments of cutaneous dermal atrophy within trials have largely been limited to clinical observation or symptom and sign reporting by trial participants (which

will only detect severe atrophy of the dermis). More sensitive and reliable methods, such as high frequency ultrasound, are routinely available (Cossmann 2006). Dermal atrophy might be monitored in part if trials routinely adopted more robust methods, such as high frequency ultrasound, to assess dermal skin atrophy skinthinning, providing useful information for patients and clinicians.

Excessive use of topical corticosteroids can also cause a substantial thinning of the epidermis (Kao 2003). However, thickening of the epidermis is part of the problem in psoriasis and so thinning of the epidermis (not dermis) that has been caused by topical corticosteroids may be of benefit in psoriasis. Evidence drawn from healthy volunteers or severe long-term cases cannot be generalised. Specifically, evidence is required concerning the frequency and spectrum of atrophy in people with chronic psoriasis requiring long-term treatment.

Topical vitamin D analogues (calcipotriol, tacalcitol) and topical calcineurin inhibitors (tacrolimus and pimecrolimus) do not cause cutaneous atrophy. The issue may become a historical one if newer safer steroids, currently being developed and evaluated for atopic dermatitis, subsequently demonstrate efficacy and safety in chronic plaque psoriasis.

We found no evidence on tachyphylaxis, but if treatment response were to decline, this could lead to over-use, increasing the risk of percutaneous absorption. As the evidence base on longer-term adverse effects in psoriasis is inadequate, the preferences of people with psoriasis and their attitudes to these perceived risks should inform treatment choice. Further research is required to inform approaches to long-term maintenance.

Implications for research

Evidence showing that treatments improve the symptoms of psoriasis has focused mainly on treatments with relatively short duration. Although improving, there is still relatively limited randomised evidence to tell us about the long-term effect of using these treatments; good quality head-to-head evidence is therefore needed to quantify and compare long-term adverse events and to explore the feasibility of long-term treatment. There are important sources of heterogeneity in currently available trial findings, which it is not possible to explore in anything other than a qualitative sense. For example, the properties of the vehicle preparation are known to deliver wide variation in response to treatment. This is important when interpreting the findings of this review. The value of the active ingredient may be worth one point on a six-point (IAGI) scale, but possibly one to two points are also being contributed by the vehicle. However, an analysis of vehicle performance was outside the scope of the review. Trial publications included in this review span 45 years. Reporting standards within these trials are generally suboptimal by today's standards, and it would be useful to ensure that current trials adhere to CON-SORT (CONsolidated Standards of Reporting Trials) standards to help future reviews to interpret findings appropriately. For example, where trials enrol participants with a wide range of baseline severity, stratifying the randomisation by baseline severity would be a useful design feature. We are not aware of studies that have adopted this approach. Trialists might usefully consider including more homogeneous participant groups in terms of severity, so that the clinical implications of findings are clearer.

Given the importance of safety to patients and to resolve clinical uncertainties, it would be valuable to obtain reliable data on the safety of corticosteroids used long-term at recommended doses, including step-down or intermittent management. Historically, these uncertainties have been driven by unrepresentative case series of non-standard use of steroids. Given the variety of methods used to assess atrophy and other sequelae, it would be valuable to reach clinical consensus about reliable assessment methods (Cossmann 2006).

ACKNOWLEDGEMENTS

The Cochrane Skin Group editorial base wishes to thank Luigi Naldi who was the Key Editor for this review; Jo Leonardi-Bee and Philippa Middleton who were the Statistical and Methods Editors, respectively; the clinical referees, Phyllis Spuls and Steven Chow; and the consumer referee, Carolyn Hughes.

The review team are indebted to Jane Harrison (formerly of the Centre for Reviews and Dissemination, University of York) for devising the original search strategies in 1999, Julie Glanville (formerly of the Centre for Reviews and Dissemination, University of York) for running the effectiveness and adverse events search strategies in 2002 and 2005, Kate Light and Kath Wright (Centre for Reviews and Dissemination, University of York) for updating the searches in 2008 and 2011, and Finola Delamere and Liz Doney of the Cochrane Skin Group for searching the Cochrane Skin Group's Specialist Skin Trials Register. We also acknowledge the valuable contribution made by Gladys Edwards, who co-authored a previous version of this review.

Last but not least, we would like to thank the authors and sponsors who provided unpublished data, which has greatly enriched this review.

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* Indicates the major publication for the study

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Agrup 1981

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 11 Treatment duration: 3 wks; FU: 3 wks LF: 0 (0%) BC: not reported Age: not reported Gender (per cent men): not reported Severity: not reported INCLUSION CRITERIA • Chronic plaque psoriasis • Stable symmetrical lesions of the same morphology • Adult EXCLUSION CRITERIA • Pregnancy • Receiving steroid preparations
Interventions	 Budesonide ointment 0.025% BD (B) Placebo (vehicle) BD (P)
Outcomes	 Investigator's preference Patient's preference
Notes	The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Topical treatments for chronic plaque psoriasis (Review)

Agrup 1981 (Continued)

Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Alora-Palli 2010

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated list Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 60 Treatment duration: 12 wks; FU: 18 wks LF: 5 (8.3%) BC: Yes Age: 48.5 (15.4SD); range = 19 to 77 Gender (per cent men): 56.7% Severity: mPASI = 7.09 (3.14SD); PGA = 3.05 Duration (yrs): 16.5 (13.0SD); range = 1 to 62 INCLUSION CRITERIA • People aged > = 18 with moderate plaque psoriasis • BSA: 3% to 15% (excluding head, groin, palms, and soles) EXCLUSION CRITERIA • Pregnancy or lactation • Topical or UVB therapy within previous 2 wks • Systemic corticosteroids, PUVA, or laser phototherapy within previous 4 wks • Other systemic therapies or biologicals within previous 12 wks
Interventions	 Calcipotriol cream 0.005% BD (C) Liquid carbonis distillate (LCD) 15% solution BD (T)
Outcomes	 Per cent change modified PASI (0 to 64.8) Physician's Global Assessment (PGA): 6-pt (0 = no disease to 5 = very severe). Overall Patient Symptom Score (erythema, thickness, burning, flaking, etc): 7-pt continuous scale (0 = none to 6 = severe) Success rates Participant satisfaction (cosmetic acceptability)

Alora-Palli 2010 (Continued)

	 6. Dermatology Quality of Life Index (DLQI) (0 to 30, where higher scores indicate poorer QoL) 7. Recurrence rates loss of PASI 50 response achieved at wk 12 PGA score at wk18 = score at wk 0
Notes	NeoStrata company, Inc. sponsored the trial. There was significant improvement in DLQI scores relative to baseline in both groups Compliance: 96% of participants in both groups reported they applied the study medi- cation twice daily on most days Recurrence rates (loss of PASI50 response): C: 7/9; T: 4/16 Recurrence rates (PGA): C: 14/20; T: 5/22 The trial author supplied unpublished data. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The study was single-blind (investigator).
Randomisation method reported	Low risk	A computer-generated list was used for ran- domisation.
Loss to follow up	Low risk	8.3%
Baseline assessments	Low risk	The trial reported demographic and clinical characteristics.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Austad 1998

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 49 Treatment duration: 6 wks; FU: 10 wks LF: 3 (6.1%) BC: yes Age: 42.4 (13.9SD; range = 18 to 68) Gender (per cent men): 63% Severity: TSS (0 to 9) = 6.4 (0.5SD; range = Duration: 15.6 (12.3SD; range = 1 to 57) INCLUSION CRITERIA • Adults • Symmetrical plaque psoriasis • Total Severity Score $\geq 6/9$ EXCLUSION CRITERIA • Widespread psoriasis • Hypercalcaemia • Liver or renal disease • Risk of pregnancy • Pregnancy • Relevant concomitant medication or con- • Previous adverse response	
Interventions	 Clobetasol propionate ointment 0.05% BD (2/52), followed by calcipotriol 50 mcg/g BD (4/52) (CP) Calcipotriol 50 mcg/g BD (6/52) (C) 	
Outcomes	 Overall severity score (0 to 9) Investigator Global Assessment (6-pt: worsened to cleared) Treatment preferences, investigator Treatment preferences, patients Compliance 	
Notes	Glaxo Wellcome Research and Developmer	nt, Norway, sponsored the trial
Risk of bias		
Bias	Authors' judgement	Support for judgement

Austad 1998 (Continued)

Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	High risk	The trial did not report this.
Loss to follow up	Low risk	6.1%
Baseline assessments	Low risk	The trial did not report these.
Baseline comparability demonstrated	Low risk	-

Baiocchi 1997

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: block randomisation (4 participants) Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Described
Participants	 N: 132 Treatment duration: 8 wks; FU: 8 wks LF: 2 (1.5%) BC: yes Age: 46.8 (15.2SD; range = 18 to 89) Gender (per cent men): 67.4% Severity: PASI = 4.4 (2.1SD) INCLUSION CRITERIA Adult Symmetrical mild-to-moderate chronic plaque psoriasis EXCLUSION CRITERIA Recent topical or systemic antipsoriatic therapy Rapidly worsening psoriasis Concurrent vitamin D Renal or hepatic disease Pregnancy Lactation
Interventions	 Calcipotriol ointment 50 mcg/g OD (C1) Calcipotriol ointment 50 mcg/g BD (C2)

Baiocchi 1997 (Continued)

Outcomes	 PASI Severity: erythema, scaling, induration (0 to 4 each) Global improvement score (7-pt: 0% to 90 - 100%) Cosmetic acceptability
Notes	The trial did not report on sponsorship. All participants had a bath with salicylic acid 3 to 4 days before starting study treatments

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Low risk	Block randomisation was used.
Loss to follow up	Low risk	1.5%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Barker 1999 (H)

Methods	DESIGN Within-patient Participant delivery Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described
Participants	N: 30 Treatment duration: 8 wks; FU: 8 wks LF: 4 (13.3%) BC: demographics similar; clinical characteristics not reported Age: 47.2 (14.5SD, N = 144) (range = 20 to 75) Gender (per cent men): 59.7% (86/144) Severity: not reported INCLUSION CRITERIA • Chronic plaque psoriasis

Barker 1999 (H) (Continued)

	 Stable bilateral lesions affecting < 20% total body surface area Adult (aged 18 to 85) EXCLUSION CRITERIA Pregnancy Concomitant disease Known hypersensitivity to vitamin D derivatives Systemic treatments within previous 1 mth Systemic retinoids within previous 2 mths Plaques < 10 cm² or > 150 cm²
Interventions	Dose-ranging study including placebo; calcipotriol 50 mcg/g; maxacalcitol 6, 12.5, 25, and 50 mcg/g OD Contrast included the following: • maxacalcitol 25 mcg/g OD • Calcipotriol 50 mcg/g OD
Outcomes	 Psoriasis Severity Index (PSI): sum scores for erythema, induration, and scaling (0 to 24) IAGI (6-pt: worse to cleared) PAGI (6-pt: worse to cleared) Investigator side preference Patient side preference
Notes	Non-target plaques received emollient or coal tar throughout. Chugai Pharma Europe sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	13.3%
Baseline assessments	Low risk	These were partially reported (demograph- ics only).
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Barker 1999 (P)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (unclear) WITHDRAWAL/DROPOUT Described
Participants	 N: 60 Treatment duration: 8 wks; FU: 8 wks LF: 6 (10.0%) BC: demographics similar; clinical characteristics not reported Age: 47.2 (14.5SD, N = 144) (range = 20 to 75) Gender (per cent men): 59.7% (86/144) Severity: not reported INCLUSION CRITERIA Chronic plaque psoriasis Stable bilateral lesions affecting < 20% total body surface area Adult (aged 18 to 85) EXCLUSION CRITERIA Pregnancy Concomitant disease Known hypersensitivity to vitamin D derivatives Systemic reatments within previous 1 mth Systemic retinoids within previous 2 mths Plaques < 10 cm² or > 150 cm²
Interventions	 Dose-ranging study including placebo; calcipotriol 50 mcg/g; maxacalcitol ointment 6, 12.5, 25, and 50 mcg/g OD Contrast included the following: Calcipotriol ointment 50 mcg/g OD (C) Placebo ointment (vehicle) (P)
Outcomes	 Psoriasis Severity Index (PSI): sum scores for erythema, induration and scaling (0 to 24) IAGI (6-pt: worse to cleared) PAGI (6-pt: worse to cleared) Investigator side preference Patient side preference
Notes	Non-target plaques received emollient or coal tar throughout the study. Chugai Pharma Europe sponsored the trial.

Risk of bias

Barker 1999 (P) (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	10.0%
Baseline assessments	Low risk	These were partially reported (demograph- ics only).
Baseline comparability demonstrated	Unclear risk	These were partially done.

Barrett 2005

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Not described
Participants	N: 420 Treatment duration: 8 wks; FU: 8 wks LF: unclear BC: not reported Age: not reported Gender (per cent men): not reported Severity: not reported INCLUSION CRITERIA • People with mild plaque psoriasis EXCLUSION CRITERIA • Not reported
Interventions	 Calcipotriol scalp 50 mcg/g solution BD, plus non-medicated shampoo twice/wk (Johnson's baby shampoo®) (CP) Calcipotriol scalp 50 mcg/g solution BD, plus tar shampoo (Polytar liquid ®) twice/wk (CT)

Barrett 2005 (Continued)

	At visit 1, all participants were treated with calcipotriol scalp solution twice daily. Par- ticipants were randomly assigned to treatment with either twice-weekly Polytar® liquic or a non-medicated shampoo twice weekly	
Outcomes	 Investigator assessment of global improvement (6-pt: worse to cleared) Total Sign Score (scale: 0 to 12) 	
Notes	Leo Pharmaceuticals sponsored the trial. The sponsor supplied unpublished outcomes data	

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial did not report this.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open (impossible to blind par- ticipants when tar-based products are used)
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Bernhard 1991 (1)

Methods	DESIGN Within-patient Delivery unclear ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 100 Treatment duration: 2 wks; FU: 2 wks LF: 4 (4%) BC: yes Age: 49 (range = 20 to 77) Gender (per cent men): 61.5%

Bernhard 1991 (1) (Continued)

	 Severity: at least 2 signs or symptoms ≥ 2 on a 4-pt scale Duration (yrs): 18.2 (range = 1 to 53) INCLUSION CRITERIA Bilateral, comparable psoriasis of at least moderate severity Adult At least 2 signs or symptoms ≥ 2 on a 4-pt scale EXCLUSION CRITERIA Not reported
Interventions	 Halobetasol 0.05% ointment BD (H) Placebo (Vehicle) (P)
Outcomes	 Signs: erythema, plaque elevation, scaling, overall lesion severity Patient Global Assessment (5-pt: poor to excellent) Skin atrophy
Notes	Westwood-Squibb Pharmaceuticals (BMS) sponsored the trial with an educational grant

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	4.0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Bernhard 1991(2)

Methods	DESIGN Between-patient Delivery unclear ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 72 Treatment duration: 2 wks; FU: 2 wks LF: 0 (0%) BC: yes (demographics); clinical comparability unclear Age: 53 (range = 23 to 86) Gender (per cent men): 52.8% Severity: signs > = 4 on a 7-pt scale; BSA = 1% to 20% Duration: 22.7 (range = 1 to 62) INCLUSION CRITERIA • Plaque psoriasis of at least moderate severity • Adult • Signs \geq 4 on a 7-pt scale • BSA 1% to 20% EXCLUSION CRITERIA • Not reported	
Interventions	 Halobetasol 0.05% ointment, BD (H) Placebo (Vehicle) (P) 	
Outcomes	 Signs: erythema, induration, scaling Investigator Global Assessment (5-pt: worse to clear) 	
Notes	Westwood-Squibb Pharmaceuticals (BMS) sponsored the trial with an educational grant	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Allocation conceannent (selection bias)	Chercal HSK	The that reported insumercint details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%

Bernhard 1991(2) (Continued)

Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.
Bernstein 2006		
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not Concealment: unclear BLINDING Double-blind (participant/inve WITHDRAWAL/DROPOUT Described	estigator)
Participants	INCLUSION CRITERIA • People aged 18 to 80 with • Good general health EXCLUSION CRITERIA • Painful/inflamed lesions • Intertriginous psoriasis • Hypertrophic lesions • Severe psoriasis • Use of topical antipsoriati • Use of systematic antipsor) 0 (2.8SD); QLI (0 to 120) = 58.74 (31.5SD) n mild to moderate plaque psoriasis (BSA < 15%) ics within previous 2 wks riatics within previous 4 wks ls, immunosuppressants, COX2s
Interventions	 <i>Mahonia aquifolium</i> (Reli Placebo (vehicle) BD (P) 	iéva™) in Novasome cream® BD (MA)
Outcomes	psoriasis involvement" (E+I+S) 2. QLI (Quality of Life Inde	an on a 4 x 4 cm section of skin "typical of the patient's) x per cent involvement (0 to 12) ex) assessed by participant. 12 questions each scored 0 Maximum score = 120 (equating to very poor quality of s.

Bernstein 2006 (Continued)

Notes	Apollo Pharmaceuticals sponsored the trial.
	'PASI' is similar to TSS (range = 0 to 12) as it examines small BSA. However, the PASI
	score was also adjusted by area

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	14.5%
Baseline assessments	Low risk	The trial reported this.
Baseline comparability demonstrated	Low risk	-

Berth Jones 1992b

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: balanced blocks of 4 using computer-generated random num-
	bers
	Concealment: unclear
	BLINDING
	Open
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 478
1 articipants	Treatment duration: 8 wks; FU: 8 wks
	LF: for PASI: 56 (11.7%); for Response: 20 (4.2%)
	BC: yes
	Age: 44 (range = 18 to 85)
	Gender (per cent men): 55%
	-
	Severity: PASI = 9.3 Duration (yrs): 18 (12SD)
	INCLUSION CRITERIA
	• Outpatients

Berth Jones 1992b (Continued)

	 Adults Chronic stable plaque psoriasis EXCLUSION CRITERIA Previous non-response to study medications Recent systemic treatment Hypercalcaemia Abnormal renal/hepatic function Calcium or vitamin D intake Relevant concomitant medication Pregnancy Risk of pregnancy
Interventions	 Calcipotriol ointment 50 mcg/g BD (C) Dithranol cream (dose titration 0.1% to 2%) OD (D)
Outcomes	 PASI Investigator Global Assessment (5-pt: worse to cleared) Patient Global Assessment (5-pt: worse to cleared) Cosmetic acceptability Compliance
Notes	Leo Pharmaceutical Products, Denmark, sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Low risk	A computer-generated block list was used for randomisation.
Loss to follow up	Low risk	11.7%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Beutner 2006

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 27 Treatment duration: 4 wks; FU: 4 wks LF: 2 (7.4%) BC: yes Age: 51.6 (12.8SD); range = 21 to 75 Gender (per cent men): 67% Ethnicity (% white): 85% Severity: overall target severity score (0 to 4) = 2.67 (0.58SD) INCLUSION CRITERIA • People aged > = 18 with moderate to severe plaque psoriasis (overall plaque severity score (0 to 8) > = 5) • 2 bilateral plaques of equivalent size (5 cm ² to 10 cm ²) EXCLUSION CRITERIA • Use of topical antipsoriatic therapy or UV exposure within previous 4 wks • Pregnancy or risk thereof
Interventions	 Clobetasol propionate 0.05% spray BD (CP) Placebo (vehicle) spray BD (P)
Outcomes	 Overall target plaque severity score using a collapsed 9-pt scale: none (0 to 1), mild (2 to 3), moderate (4 to 5), severe (6 to 7), and very severe (8) Signs: scaling, erythema, and plaque elevation (each scored 0 to 8) Adverse events (burning, stinging, pruritus, telangiectasias, skin atrophy)
Notes	Galderma Laboratories, L.P. sponsored the study. There was SD imputation (TSS).

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Beutner 2006 (Continued)

Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	7.4%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Bourke 1993b

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: participants were randomised into groups A and B, then randomised to left/right application with sealed envelopes Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 19 (evaluable) Treatment duration: 8 wks; FU: 8 wks LF: NR BC: yes (clinical only) Age: not reported Gender (per cent men): not reported Severity: TSS = 7.9 INCLUSION CRITERIA • Adult • Symmetrical chronic plaque psoriasis • Outpatients EXCLUSION CRITERIA • UV or systemic antipsoriatic therapy
Interventions	 Calcipotriol BD (C) Calcipotriol BD plus polythene film at night (O)
Outcomes	 Signs: erythema, induration, scale Total Sign Score (0 to 12)
Notes	Participants were randomised into groups A (calcipotriol BD) and B (occlusion ON), then each participant was randomised to left/right application: group A (occlusion ON/ no occlusion); group B (calcipotriol BD or placebo BD). The study reported findings for group A. The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was single-blind (investigator).
Randomisation method reported	Low risk	Envelopes were used.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Bourke 1997

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described
Participants	N: 24 Treatment duration: 8 wks LF: 4 (16.7%) BC: yes (clinical only reported) Age: not reported Gender (per cent men): 41.7% Severity: PASI mean = 14.0 INCLUSION CRITERIA • Adults • Symmetrical chronic moderate chronic plaque psoriasis EXCLUSION CRITERIA • Pregnancy • Lactation • Drugs affecting systemic calcium homeostasis • Recent systemic antipsoriatic or UVB therapy

Bourke 1997 (Continued)

Interventions	 Calcitriol 3 mcg/g BD (CL) Calcipotriol 50 mcg/g BD (C)
Outcomes	1. PASI
Notes	Solvay-Duphar Ltd sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	16.7%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Brown 2005

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not stated
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 30
rancipants	
	Treatment duration: 12 wks; FU: 12 wks
	LF: 6 (20%)
	BC: unclear
	Age: 55 (12.6SD); range = 20 to 75
	Gender (per cent men): 50%
	Ethnicity (% white): 46.7%
	Severity: TSS (0 to 16) mean = 6.97
	INCLUSION CRITERIA

	 People with mild stable plaque psoriasis BSA affected < 15% General good health EXCLUSION CRITERIA Pregnancy or risk thereof Lactation Phototherapy, topical therapy, or systemic therapy within previous 4 wks Cancer History of drug or alcohol abuse Concomitant antipsoriatic therapy
Interventions	 Kukui nut oil TD (K) Placebo (mineral oil, vehicle) TD (P)
Outcomes	 Investigator assessment of PASI (scale unclear (0 to 64.8)) and Global Severity Score (5-pt: 0 = none to 4 = very severe) Subject assessment of Global Severity Score (5-pt: 0 = none to 4 = very severe) Total Severity Score (0 to 16; thickness + scaliness + erythema + itch) Compliance also assessed (bottles weighed)
Notes	Hawaii Community Foundation sponsored the study. The trial author supplied unpublished data. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	20%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Bruce 1994

· ·			
Methods	Concealment: unclear BLINDING	Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT	
Participants	LF: 15 (13.2%) BC: yes Age: 44.1 (14.6SD; range = 20 to 77) Gender (per cent men): 60.2% Severity: mean duration of current epi Overall severity score (mean): 4.5 INCLUSION CRITERIA • Stable plaque psoriasis • Adults • At least mild overall severity • At least moderately severe plaque EXCLUSION CRITERIA • Pregnancy • Lactation • Inadequate contraception • Sensitivity to test medications • Recent topical, UV, or systemic to	N: 114 Treatment duration: 6 wks; FU: 6 wks LF: 15 (13.2%) BC: yes Age: 44.1 (14.6SD; range = 20 to 77) Gender (per cent men): 60.2% Severity: mean duration of current episode (days) = 142 (range = 0 to 601) Overall severity score (mean): 4.5 INCLUSION CRITERIA • Stable plaque psoriasis • Adults • At least mild overall severity • At least moderately severe plaque elevation EXCLUSION CRITERIA • Pregnancy • Lactation • Inadequate contraception • Sensitivity to test medications • Recent topical, UV, or systemic treatment • Recent involvement in other trials	
Interventions		 Calcipotriol ointment 0.005% BD (C) Fluocinonide ointment 0.05% BD (F) 	
Outcomes		 Sign: scaling, erythema, plaque elevation Overall severity (Total Sign Score and per cent involvement) Investigator Global Assessment 	
Notes	Westwood Squibb Pharmaceuticals In There was SD imputation (TSS).	Westwood Squibb Pharmaceuticals Inc. sponsored the trial. There was SD imputation (TSS).	
Risk of bias			
Bias	Authors' judgement	Support for judgement	

Allocation concealment (selection bias) Unclear risk The trial reported insufficient details.

Bruce 1994 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	13.2%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Buckley 1978

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 10 Treatment duration: 3 wks; FU: 3 wks LF: 2 (20%) BC: not reported Age: 21.4 (range = 9 to 41) Gender (per cent men): 50% Severity: not reported INCLUSION CRITERIA • Active chronic psoriasis • Lesions approximately symmetrically distributed EXCLUSION CRITERIA • Not reported
Interventions	 Dead Sea salts emollient lotion 30% (frequency of application not reported) (D) Base emollient lotion (placebo) (P)
Outcomes	 Jacoby assessment score (0 to 7 score transformed to per cent clinical improvement) Photographic evaluation Overall patient assessment (relative efficacy, speed of response, irritation, staining, ease of application)

Buckley 1978 (Continued)

Notes	The trial did not report sponsorship.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.	
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).	
Randomisation method reported	Unclear risk	The trial did not report this.	
Loss to follow up	Low risk	20.0%	
Baseline assessments	Low risk	These were partially done.	
Baseline comparability demonstrated	Unclear risk	The trial did not report this.	

Buckley 2008

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator)	
Participants	WITHDRAWAL/DROPOUT Described N: 218 Treatment duration: 8 wks; FU: 10 wks	
	Treatment duration: 8 wks; FU: 10 wks LF: 5 (2.3%) BC: yes Age: 48.4 (15.48SD) Gender (per cent men): 45.0% Ethnicity: 97.7% Severity: TSS (0 to 12) = 6.80 (1.58SD) Duration (yrs): 14.6 (13.87SD); range = 0 to 65 Extent of scalp psoriasis: 3.39 (1.36SD) INCLUSION CRITERIA • People aged > = 18 with scalp psoriasis affecting > = 10% scalp • Amenable to topical treatment with < = 100 g medication/wk	

Buckley 2008 (Continued)

	 TSS (0 to 12) > = 4 Each individual sign score (0 to 4) > = 1 IGA at least mild (> = 3) EXCLUSION CRITERIA Erythrodermic psoriasis Pustular psoriasis Systemic or PUVA therapy within previous 4 wks UVB or grenz ray therapy on scalp, or topical scalp therapy within previous 2 wks Severe renal impairment or severe hepatic disorders
Interventions	 Combined gel: calcipotriol 50 mcg/g + betamethasone dipropionate 0.5 mg/g OD (C-B) Betamethasone dipropionate 0.5 mg/g gel OD (B) Participants achieving absence of disease (IGA = 5) at weeks 2 to 5 could withdraw from the study
Outcomes	 Signs: redness, thickness, scaliness (each sign scored on 5 pt scale 0 to 4) Total Sign Score (TSS; 13-pt: 0 = none to 12 = very severe symptoms) Investigator's Global Assessment (IGA): 0 = absence of disease to 5 = very severe disease. Controlled disease: IGA < = 1 Investigator assessment of extent of scalp psoriasis: 0% to 100% (score of 3 corresponds to 30% to 49% involvement) Patient's assessment of global improvement (PAGI): 7-pt scale: 0 = worse to 6 = cleared. Reported as dichotomised treatment success: PAGI > = 4 Treatment duration Compliance (medication usage)
Notes	Leo Pharma A/S, Ballerup, Denmark, sponsored the study. Treatment duration (wks): C-B: 6.1 (2.4 SD), N = 108; B: 6.8 (2.2 SD), N = 110 Compliance (missed < = 20% applications): C-B: 93/108; B: 103/110 The sponsor supplied unpublished data.,

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	2.3%
Baseline assessments	Low risk	These were reported.

Buckley 2008 (Continued)

Baseline comparability demonstrated	Low risk -
Camarasa 2003	
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Conducted in 20 centres; stratification not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 258 Treatment duration: 6 wks; FU: 14 wks LF: 15 (5.8%) BC: yes Age: 43.5 (14.3SD: range = 15 to 83) Gender (per cent men): 64.3% Severity: per cent BSA = 25.5 (22.9SD: range = 1 to 95); PASI = 15.4 (10.6SD) Duration of psoriasis (mths) mean: 199.2 (157.5SD: range = 1 to 745) INCLUSION CRITERIA Adults Moderate to severe chronic plaque psoriasis (≥ 2 on global severity score) EXCLUSION CRITERIA Systemic or intralesional therapy or photo(chemo)therapy in previous 2 mths Medications or conditions that might interfere with the assessment of study drugs Concomitant bacterial, fungal, or viral skin conditions Clinically relevant abnormalities in laboratory parameters (calcium homeostasis and renal function) Pregnancy or lactation Absence of adequate contraception, where appropriate
Interventions	 Calcitriol 3 mcg/g ointment BD (C) Betamethasone dipropionate 0.05% ointment BD (B)
Outcomes	 IAGI (6-pt: worsening to clearance) PASI Overall global severity of lesions (5-pt: 0 = none to 4 = very severe) Relapse rate Proportion remaining in remission (non-randomised subgroup analysis)

Camarasa 2003 (Continued)

Notes	There was a 1-wk run-in period without treatment, except tar shampoo and emollients. Follow up was for responders only (defined as achieving clearance or considerable im-
	provement). The scalp was excluded.
	Galderma Laboratories sponsored the trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	5.8%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Cheesbrough 1992

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 24
Tarticipants	Treatment duration: 12 wks; FU: 12 wks
	LF: 5 (20.8 %)
	BC: yes
	Age (mean): 47
	Gender (per cent men): 54.2%
	Severity: PASI mean = 26
	INCLUSION CRITERIA
	Chronic stable plaque psoriasis
	· Ontoine stable plaque postasis

Cheesbrough 1992 (Continued)

	EXCLUSION CRITERIANot reported
Interventions	 Dead sea salts emollient lotion 30% (frequency of application not reported) (D) Base emollient lotion (placebo) (P)
Outcomes	 PASI Erythema, scaling, thickening, pruritus Adverse events
Notes	Finders Dead Sea Salt Co and Dead Sea Salt works supplied the study treatments

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	20.8%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Christensen 1999

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Single-blind at inclusion only (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 171 Treatment duration: 8 wks; FU: 16 wks (N = 95) LF: 5 (2.9%) BC: yes

Christensen 1999 (Continued)

	Age: 47.4 (range = 17 to 88) Gender (per cent men): 62.6% Severity: mean TSS (0 to 9) = 6.24; mean duration of psoriasis = 18.5 (range = 1 to 58) INCLUSION CRITERIA • Outpatients with mild to severe chronic stable chronic plaque psoriasis, not more than 10% BSA, Total Severity Score (0 to 9) \geq 4, involving all 3 signs (erythema, scaling, infiltration) EXCLUSION CRITERIA • Systemic treatment within previous 4 wks • Topical treatment within previous 2 wks; receipt of oral retinoids within previous 2 mths
Interventions	 Short-contact dithranol (30 min) 1% to 3%, OD (D) Calcipotriol 50 mcg/g BD (C)
Outcomes	 Total Severity Score (0 to 9) Pruritus (0 to 3) Investigator's Global Assessment (7-pt: worse to completely clear) Patient's Global Assessment (6-pt: worse to completely clear) Adverse events
Notes	The study did not report sponsorship. The study also included 16-week follow-up data for consenting participants who achieved at least 50% improvement from baseline (Investigator scale). There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	2.9%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Cook-Bolden 2010

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 81 Treatment duration: 4 wks; FU: 4 wks LF: 3 (4%) BC: yes Age: 43.7 (14.69SD) Ethnicity (% white): 78% Gender (per cent men): 39.5% Duration (yrs): 11.1 (10.5SD) Severity: GSS = 3 (moderate) (67.9%); GSS = 4 (severe) (32.1%) INCLUSION CRITERIA • People aged 18 or over with moderate to severe plaque psoriasis of the scalp (GSS = 3 or GSS = 4) EXCLUSION CRITERIA • Use of chemical hair process, steroid medication, ultraviolet B (UVB) treatment, or both; calcipotriene; other vitamin D analogues; anthralin/tar; all other anti-psoriasis medications (previous 2 wks) • Use of PUVA or other non-biological systemic treatments (previous 4 wks) • Use of biological therapies (previous 12 wks) • Pregnancy or risk thereof • Lactation
Interventions	 Clobetasol propionate spray 0.05% BD (CP) Placebo (vehicle) spray BD (P) Max usage/wk: 50 g Participants achieving GSS = 0 at 2 wks completed ("withdrew" from) the study
Outcomes	 Global Severity Score (GSS) (6-pt: 0 = none to 5 = very severe), dichotomised as success (clear/almost clear) and failure Scalp psoriasis sign scores: erythema, scaling, elevation (0 to 4, where 0 = none) Pruritus (4-pt: 0 = no itching to 3 = intense itching that disrupts sleep) Atrophy (0 to 3, where 0 = none) Telangiectasias (0 to 3, where 0 = none) Stinging/burning (0 to 3, where 0 = none) Patient satisfaction with applicator Scalpdex, a scalp dermatitis-specific quality of life instrument; 23 questions, each scored 0 (never) to 100 (all the time) Compliance: yes if participant achieved 80% to 120% of expected applications

Cook-Bolden 2010 (Continued)

Notes	Galderma Laboratories, LP, sponsored the trial.
	Galderma supplied safety data for this study.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	4%
Baseline assessments	Low risk	These were made.
Baseline comparability demonstrated	Low risk	The trial demonstrated demographic and clinical comparability

Crosti 1997

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Unclear
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 160
	Treatment duration: 6 wks; FU: 10 wks
	LF: 8 (5%)
	BC: yes
	Age: 49.9 (14.2SD)
	Gender (per cent men): 68.1%
	Severity: mean PASI = 7.6
	INCLUSION CRITERIA
	Mild stable chronic plaque psoriasis
	• Adult
	EXCLUSION CRITERIA
	• Recent topical or systemic treatments

Crosti 1997 (Continued)

	 Pregnancy Lactation Concomitant vitamin D or systemic steroids Hepatic or renal failure
Interventions	 Calcipotriol ointment 50 mcg/g BD (C) Betamethasone dipropionate + salicylic acid BD (B)
Outcomes	 PASI Investigator Global Assessment Patient Global Assessment of acceptability of treatment (5-pt: nil to excellent)
Notes	The trial did not report sponsorship. There was SD imputation (PASI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	5.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Cunliffe 1992

Methods	DESIGN Between-patient Delivery unclear ALLOCATION Random Method of randomisation: balanced blocks of 10 according to a computer-generated random numbers table Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 409 Treatment duration: 6 wks; FU: 6 wks LF: 8 (2.0%) BC: yes Age: 44.9 (range = 17 to 83) Gender (per cent men): 55.7% Severity: mean PASI = 9.0 (range = 0.6 to 41.2); mean duration psoriasis = 16.2 (range = 0.2 to 57) INCLUSION CRITERIA • Stable plaque psoriasis • Adults • Outpatients EXCLUSION CRITERIA • Risk of pregnancy • Pregnancy • Lactation • Recent systemic antipsoriatic treatment
Interventions	 Calcipotriol ointment 50 mcg/g BD (C) Betamethasone 17-valerate 1 mg/g BD (B)
Outcomes	 PASI Patient overall assessment (5 pt: worse to clear)
Notes	Leo Pharmaceuticals sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Cunliffe 1992 (Continued)

Randomisation method reported	Low risk	A computer-generated block list was used for randomisation.
Loss to follow up	Low risk	2.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

De Simone 1993

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Blinding unclear WITHDRAWAL/DROPOUT Not described
Participants	N: 30 Treatment duration: 6 wks; FU: 10 wks LF: 0 (0%) BC: not reported Age: range = 18 to 84 Gender (per cent men): 70.0% Severity: PASI range = 2.7 to 24.3 INCLUSION CRITERIA • Chronic plaque psoriasis EXCLUSION CRITERIA • Pregnancy • Lactation • Hepatic or renal disease • Recent systemic or topical therapy • High intake of vitamin D or calcium
Interventions	 Calcipotriol ointment 50 mcg/g BD (C) Coal tar 5% in Lassar's paste (T)
Outcomes	1. Investigator Global Assessment (estimated from PASI score)
Notes	The trial did not report sponsorship.

De Simone 1993 (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Decroix 2004

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Single-blind (investigator); as cream and lotion were compared, not possible to blind participants WITHDRAWAL/DROPOUT Described
Participants	N: 222 Treatment duration: 4 wks; FU: 4 wks LF: 0 (0%) BC: yes Age: 48.4 (15.0SD) Gender (per cent men): 55.9% Ethnicity (% white): 100% Severity: DSS (0 to 12) = 8.44 (1.45SD); atrophy (0 to 3) = 1.01; telangiectasia (0 to 3) = 0.01 INCLUSION CRITERIA • People aged > = 18 with stable moderate to severe plaque psoriasis • Target lesion diameter > 3 cm • Lesion not localised to scalp, face, hands, or feet • BSA > = 10%; • (Women participants only) negative pregnancy test EXCLUSION CRITERIA • Use of topical treatments and phototherapy within previous 2 wks

Decroix 2004 (Continued)

	Use of systemic therapies within previous 2 to 16 wksRegular sun exposure within previous 2 wks
Interventions	 Clobetasol propionate lotion/cream BD (CP) Placebo (vehicle lotion) BD (P) Participants were randomised to clobetasol propionate cream or lotion; results were aggregated for review purposes
Outcomes	 Erythema, plaque elevation, scaling pruritus: each scored on 5-pt scale (0 to 4) Dermatological Sum Score (DSS: erythema, elevation, scaling): 0 to 12 Global severity (GSS): 0 (none) to 4 (severe) Dichotomised as success (GSS = 0, 0.5, or 1) or failure (GSS = 2 to 4) Investigator's Assessment of Global Improvement (IAGI): -1 (worse) to 5 (clear) Per cent BSA involvement: rule of nines Safety: telangiectasia (0 to 3), atrophy (0 to 3); adverse events Compliance (assessment method not reported) Cosmetic acceptability (questionnaire survey)
Notes	Galderma R&D, Sophia Antipolis, France, sponsored the trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator); as the cream and lotion were compared, it was not possible to blind participants
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Douglas 2002

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation schedule Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 1106 Treatment duration: 4 wks; FU: 4 wks LF: 86 (7.8%) BC: yes Age: mean = 47.1 (range = 18 to 89) Gender (per cent men): 59.8% Severity: PASI = 10.7 (range = 2.1 to 39.6) Duration: mean = 18.4 (range = 0 to 65) INCLUSION CRITERIA Chronic plaque psoriasis Aged at least 18 years Use of systemic antipsoriatic treatment/phototherapy in previous 6 weeks Treatment of lesions contraindicated for topical corticosteroid therapy EXCLUSION CRITERIA Pregnancy Lactation Current participation in other trial Abnormality of calcium metabolism Hypercalcaemia
Interventions	 Calcipotriol (50 mcg/g)/betamethasone (0.5 mg/g) combination ointment (Daviobet®) BD (D) Calcipotriol ointment (Daivonex®) 50 mcg/g BD (C) Betamethasone dipropionate ointment (Diprosone®) 0.5 mg/g BD (B) All groups then received 4 weeks of maintenance therapy with calcipotriol BD
Outcomes	 PASI (modified) (0 to 64.8) Redness, thickness, scaling (0 to 8 each) Investigator Global Assessment (6-pt: worse to cleared) Patient's assessment of treatment response (6-pt: worse to cleared) Adverse events
Notes	Leo Pharmaceuticals sponsored the trial. 4-week follow-up study also reported (open design: all participants received calcipotriol)

Risk of bias

Douglas 2002 (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	7.8%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Dubertret 1992

Methods	DESIGN
	Within-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: unclear
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 66
rancipants	Treatment duration: 4 wks; FU: 8 wks
	LF: 6 (9.1%)
	BC: yes
	Age: 43 (range = 21 to 84)
	Gender (per cent men): 69.7%
	Severity: PASI mean = 14.15
	Duration: 13.3 (range = 0.3 to 40.0)
	INCLUSION CRITERIA
	Bilateral stable symmetric chronic plaque psoriasis of the arms, limbs, or trunk
	• Adult
	EXCLUSION CRITERIA
	• Guttate or pustular psoriasis
	• Psoriasis restricted to the scalp, face, elbows, or knees
	• Recent systemic or UV therapy in the previous 10 weeks
	Calcium, vitamin D daily or other medications
	Hepatic or renal impairment
	• Pplanned exposure to sun

Dubertret 1992 (Continued)

Interventions	 Calcipotriol ointment 50 mcg/gm BD (C) Placebo (vehicle) (P)
Outcomes	 Severity [erythema, infiltration, desquamation] Modified PASI Preferred treatment Investigator Global Assessment (5-pt: cleared to worse) Patient Global Assessment (5-pt: cleared to worse)
Notes	Leo Pharmaceuticals sponsored the trial. We contacted Leo for patient outcome data. There was the SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	9.1%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Durakovic 2001

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 15 Treatment duration: 12 wks; FU: 20 wks LF: 0 (0%)

Durakovic 2001 (Continued)

	BC: yes Age: 49 (range = 27 to 76) Gender (per cent men): 80% Severity: TSS (0 to 24) = 13.7 (14.7SD) INCLUSION CRITERIA • Chronic plaque psoriasis involving at least 5% BSA • Bilateral lesions of approximately 25 cm ² EXCLUSION CRITERIA • Systemic treatment within previous 30 days • Topical treatment within previous 1 day • History of hepatic or renal failure • Nephrolithiasis; hypercalcaemia • Hypercalciuria • Pregnancy • Lactation • Unwillingness in women to use effective contraception
Interventions	 Hexafluoro-1,25-dihydroxyvitamin D 5 mcg/g in 0.1g of ointment BD (F6) Placebo ointment BD (P)
Outcomes	 Total Severity Score (0 to 24) PASI (0 to 72) Investigator's assessment of global improvement (5-pt: worsening to excellent improvement)
Notes	National Institutes of Health and by Penederm Inc. sponsored the trial. 8-week follow up (open design: all participants received study drug) study also reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Durakovic 2004

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not repo Concealment: unclear BLINDING Double-blind (participant/investiga WITHDRAWAL/DROPOUT Described		
Participants	EXCLUSION CRITERIA Previous topical therapy within Systemic therapy within previo Pregnancy or risk thereof Lactation Hepatic failure Renal failure Hypercalcaemia Hypercalciuria Hyperphosphataemia	th diameter ≥ 5cm each of plaque elevation, scaling, and erythema n previous 2 wks	
Interventions	 Paricalcitol (19-nor-1 alpha,25) ointment 15 mcg/g OD (PC) Placebo ointment OD (P) 		
Outcomes		 Global severity score (0 to 12) (erythema, plaque elevation, scaling) Global treatment success rates (IAGI) (4-pt: excellent, moderate, mild, or no improvement) 	
Notes		The trial was sponsored in part by grant from the National Institutes for Health. US Abbott Laboratories supplied the study drug	
Risk of bias			
Bias	Authors' judgement	Support for judgement	

Durakovic 2004 (Continued)

Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Duweb 2000

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Unclear WITHDRAWAL/DROPOUT Described
Participants	N: 42 Treatment duration: 6 wks; FU: unclear LF: 0 (0%) BC: clinical only Age: 33.5 (range = 6 to 61) Gender (per cent men): 69% Severity: TSS (0 to 12) = 5.2 INCLUSION CRITERIA • Psoriasis of the scalp EXCLUSION CRITERIA • Not reported
Interventions	 Calcipotriol 50 mcg/g/ml solution BD (C) Betamethasone valerate 1% lotion BD (B)
Outcomes	 Redness, thickness, scaliness (0 to 4) Total Severity Score (0 to 12) Adverse events

Duweb 2000 (Continued)

Notes	Leo Pharmaceuticals sponsored the trial. There was SD imputation (TSS).	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	This was partially demonstrated.

Elie 1983

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: unclear
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 40 (55% psoriasis)
	Treatment duration: 3 wks; FU: 3 wks
	LF: not reported
	BC: yes
	Age: 36.5 (range = 20 to 63)
	Gender (per cent men): 40%
	Severity: not reported
	INCLUSION CRITERIA
	• Moderate to severe psoriasis, seborrhoeic dermatitis, or neurodermatitis of the
	scalp
	• Adult
	EXCLUSION CRITERIA
	None reported

Elie 1983 (Continued)

Interventions	 Betamethasone-17,21-dipropionate, 0.05% BD (B) Salicylic acid 2% BD (S) Betamethasone-17,21-dipropionate, 0.05% + Salicylic acid 2% BD (BS) Placebo (vehicle) (P)
Outcomes	 Investigator Global Assessment (5-pt: very severe to clear) Severity (redness; scaling; pruritis) Area of lesion (cm²)
Notes	Schering Canada Inc. sponsored the trial. This was a scalp trial. There was SD imputation (TSS/IAGI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Ellis 1988

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated randomisation list (1:1)
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 165 Treatment duration: 3 wks; FU: 3 wks LF: 33 (20%)

	BC: yes Age: 49.1 (range = 19 to 82) Gender (per cent men): 51.6% Severity: not reported INCLUSION CRITERIA • Psoriasis of the scalp • Adult • TSS (0 to 12) \geq 6 • Particpants required to have psoriatic lesions elsewhere EXCLUSION CRITERIA • Acute systemic illness • Active skin infection • Concomitant antihistamine, topical, or systemic corticosteroid, antimetabolites, PUVA, or other dermatological treatment • Recalcitrant psoriasis • Intolerance or hypersensitivity to topical corticosteroids
Interventions	 Amerinonide lotion 0.1% OD (A) Placebo (vehicle) (P)
Outcomes	 Severity (erythema; excoriation; scaling; induration, pruritis) Total Sign Score (erythema; scaling; induration, pruritis) Investigator's Overall Evaluation (7-pt: cleared to exacerbation) Patient's Overall Evaluation (4-pt: poor to excellent) Patient Acceptability Evaluation
Notes	The trial did not report sponsorship. Compliance was checked by counting returned bottles. This was a scalp trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	20.0%
Baseline assessments	Low risk	-

Baseline comparability demonstrated	Low risk -	
Escobar 1992		
Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: 'randomised code' Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 25 Treatment duration: 4 wks; FU: 8 wks LF: 0 (0%) BC: yes Age: 40.3 (14.1SD); range = 18 to 66 Gender (per cent men): 56.0% Severity: mean TSS (0 to 12) = 7.83 INCLUSION CRITERIA • Clinical and histopathological diagnosis of psoriasis EXCLUSION CRITERIA • Systemic cytostatic/corticosteroid therapy within previous year • Renal, hepatic, haematological disease • NSAIDs, beta adrenergic receptor blockers, antimalarial drugs	
Interventions	Fish oil plus 6-hour occlusion OD (FO)Liquid paraffin plus 6-hour occlusion OD (LP)	
Outcomes	 Erythema, scaling, thickening (0 to 4) Pruritis (VAS) Patient acceptability 	
Notes	The trial did not report sponsorship. There was SD imputation (TSS).	
Risk of bias		

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Escobar 1992 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Farkas 1999

Methods	DESIGN Between-patient Delivery unclear ALLOCATION Random Method of randomisation: computer programme randomised participants in blocks of 10 Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Described
Participants	N: 84 Treatment duration: 8 wks; FU: 12 wks LF: 0 (0%) BC: yes Age: 45.1 (range = 18 to 69) Gender (per cent men): 60.7% Severity: mean PASI = 13.2%; BSA (mean) = 16.5% INCLUSION CRITERIA • Chronic stable plaque psoriasis • Adults • White participants = 30% BSA • mPASI > 10 • In- and outpatients EXCLUSION CRITERIA • Recent topical, systemic, or UV therapies • Sensitivity to study medications • Concurrent medication • Abnormal hepatic or renal function • Risk of pregnancy • Pregnancy • Lactation

	Serious co-morbidity
Interventions	 Tacalcitol ointment 4 mcg/g OD (T) Dithranol stick 1.5% or 3% OD (D)
Outcomes	 PASI (modified to exclude head) Total Sign Score (erythema, infiltration, and desquamation) Investigator Global Assessment Patients evaluation of benefit (10-pt) Investigator evaluation of efficacy and tolerability (1 = very good to 4 = very bad) Patient evaluation of efficacy and tolerability (1 = very good to 4 = very bad)
Notes	Hermal sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Low risk	A computer-generated block list was used for randomisation.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Feldman 2010 (1)

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not stated
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described

Feldman 2010 (1) (Continued)

Participants	N: 323 Treatment duration: 8 wks; FU: 8 wks LF: 7 (2.2%) BC: yes Age: 47.8 (15.2SD) Gender (per cent men): 48.3% Per cent white: 90.7% Severity: ISGA = mild disease: 26.9%; ISGA = moderate disease: 73.1%; SGA = 3.69 (0.89SD); % BSA = 6.4% (4.9SD) INCLUSION CRITERIA • People aged \geq 12 with mild to moderate plaque psoriasis • BSA: 2% to 20% • ISGA: 2 to 3 EXCLUSION CRITERIA • Known allergy to any component of formulation • Hypercalcaemia or history thereof • Pregnancy or risk thereof • Topical or systemic therapy within previous 4 wks • Previous participation in trial of study medication
Interventions	 Calcipotriol foam 0.005% BD (C) Placebo foam BD (P)
Outcomes	 Investigator Static Global Assessment (IGSA): scale unclear Treatment success: ISGA < = 1 (clear/almost clear) and minimum improvement from baseline of 2 points. Subject Global Assessment (SGA) (6-pt: 0 = clear to 5 = severe) Signs: erythema, scaling, thickness (0 to 4)
Notes	STUDY 1: nCT00689481 Stiefel Laboratories, a GSK Company, sponsored the trial. Atrophy was not reported. We sought data, but they were not received. No usable effectiveness data were reported or available from sponsor

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	2.2%

Feldman 2010 (1) (Continued)

Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Feldman 2010 (2)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 336 Treatment duration: 8 wks; FU: 8 wks LF: 6 (1.8%) BC: yes Age: 48.7 (14.4SD) Gender (per cent men): 60.4% % white: 86.0% Severity: ISGA = mild disease: 31.8%; ISGA = moderate disease: 68.2%; SGA = 3.69 (0.83SD); % BSA = 6.2% (4.8SD) INCLUSION CRITERIA • People aged \geq 12 with mild to moderate plaque psoriasis • BSA: 2% to 20% • ISGA: 2 to 3 EXCLUSION CRITERIA • Known allergy to any component of formulation • Hypercalcaemia or history thereof • Pregnancy or risk thereof • Topical or systemic therapy within previous 4 wks • Previous participation in trial of study medication
Interventions	 Calcipotriol foam 0.005% BD (C) Placebo foam BD (P)
Outcomes	 Investigator Static Global Assessment (IGSA): scale unclear Treatment success: ISGA < = 1 (clear/almost clear) and minimum improvement from baseline of 2 points Subject Global Assessment (SGA) (6-pt: 0 = clear to 5 = severe) Signs: erythema, scaling, thickness (0 to 4)

Feldman 2010 (2) (Continued)

Notes	STUDY 2: nCT00688519
	Stiefel Laboratories, a GSK Company, sponsored the trial.
	Atrophy was not reported.
	We sought data, but they were not received. No usable effectiveness data were reported
	or available from sponsor

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	1.8%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Fleming 2010 (H)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated schedule Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT
Participants	Described N: 364 Treatment duration: 8 wks; FU: 10 wks LF: 2 (0.5%) BC: yes Age: 51.1 (14.7SD) Gender (per cent men): 58.8% Per cent white: 98.1% Severity: PASI (0 to 64.8) = 7.8 (4.4SD), range = 1 to 25; IGA = 2.97 (0.66SD), range = mild (2) to very severe (5)

	Duration (yrs): 18.9 (13.8SD) INCLUSION CRITERIA • People aged > = 18 with psoriasis vulgaris affecting the trunk and or limbs, amenable to treatment with < = 100 g topical medication/w • IGA at least mild EXCLUSION CRITERIA • Guttate, erythrodermic, exfoliative or pustular psoriasis • Use of biological therapy within previous 6 mths
	 Use of systemic antipsoriatic therapy, PUVA, or grenz ray therapy with previous 4 wks Topical or UVB therapy within previous 2 wks Concomitant use of emollients during study
Interventions	 Calcipotriol gel 50 mcg/g plus betamethasone dipropionate gel 0.5 mg/g OD (C-B) Calcipotriol gel 50 mcg/g OD (C) Betamethasone dipropionate gel 0.5 mg/g OD (B) Concomitant use of corticosteroids (WHO group I or II), dithranol, tar, and retinoids on face, scalp, and flexures was permitted
Outcomes	 Investigator's Global Assessment (IGA) of disease severity; 6-pt (clear, minimal disease, mild, moderate, severe, very severe); rescaled as clear (0) to very severe (5) Responder (IGA): If IGA moderate at wk 0: clear/minimal at wk 4 or 8. If IGA mild at wk 0: clear at wk 4 or 8 PASI (modified) (0 to 64.8): change from baseline Treatment success: PASI75 Adverse events Compliance
Notes	Leo Pharma, Ballerup, Denmark, sponsored the trial. Compliance: mean weekly us =: C-B: 22.7 g; C: 22.4g; B: 25.9 g Most participants (73.5% to 80%) were fully compliant and those non-compliant missed < 10% applications Safety data were available for 362/364 participants. Leo Pharmaceuticals supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	A computer-generated randomisation schedule was used.

Fleming 2010 (H) (Continued)

Loss to follow up	Low risk	0.5%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Fleming 2010 (P)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated schedule Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 364 Treatment duration: 8 wks; FU: 10 wks LF: 2 (0.5%) BC: yes Age: 51.1 (14.7SD) Gender (per cent men): 58.8% Per cent white: 98.1% Severity: PASI (0 to 64.8) = 7.8 (4.4SD), range = 1 to 25; IGA = 2.97 (0.66SD), range = mild (2) to very severe (5) Duration (yrs): 18.9 (13.8SD) INCLUSION CRITERIA • People aged > = 18 with psoriasis vulgaris affecting the trunk and or limbs, amenable to treatment with < = 100 g topical medication/w • IGA at least mild EXCLUSION CRITERIA • Guttate, erythrodermic, exfoliative or pustular psoriasis • Use of biological therapy within previous 6 mths • Use of systemic antipsoriatic therapy, PUVA, or grenz ray therapy with previous 4 wks • Topical or UVB therapy within previous 2 wks • Concomitant use of emollients during study
Interventions	 Calcipotriol gel 50 mcg/g plus betamethasone dipropionate gel 0.5 mg/g OD (C-B) Calcipotriol gel 50 mcg/g OD (C) Betamethasone dipropionate gel 0.5 mg/g OD (B) Placebo gel OD (P) Concomitant use of corticosteroids (WHO group I or II), dithranol, tar, and retinoids

Fleming 2010 (P) (Continued)

	on face, scalp, and flexures were permitted
Outcomes	 Investigator's Global Assessment (IGA) of disease severity; 6-pt (clear, minimal disease, mild, moderate, severe, very severe); rescaled as clear (0) to very severe (5) Responder (IGA): If IGA moderate at wk 0: clear/minimal at wk 4 or 8. If IGA mild at wk 0: clear at wk 4 or 8 PASI (modified) (0 to 64.8): change from baseline Treatment success: PASI75 Adverse events Compliance
Notes	Leo Pharma, Ballerup, Denmark, sponsored the trial. Compliance: mean weekly us =: C-B: 22.7 g; C: 22.4g; B: 25.9 g; P: 26.1 g Most participants (73.5% to 80%) were fully compliant and those non-compliant missed < 10% applications Safety data were available for 362/364 participants. Leo Pharmaceuticals supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	A computer-generated randomisation schedule was used.
Loss to follow up	Low risk	0.5%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Franz 1999

Methods	DESIGN Between-patient Participant delivery ALLOCATION	
	Random Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT	
	Described	
Participants	N: 190 Treatment duration: 2 wks; FU: 4 wks LF: 18 (9.5%) BC: yes Age: 49.6 Gender (per cent men): 49.3% Severity: mean TSS (0 to 12) = 7.92 INCLUSION CRITERIA • Moderate to severe scalp psoriasis (each of 3 primary signs ≥ 2) • Scalp involvement $\geq 10\%$ • Adults EXCLUSION CRITERIA • Systemic psoriatic therapy within previous 4 wks • Topical scalp preparations within previous 2 wks	
Interventions	 Betamethasone valerate foam 0.1% BD Placebo foam BD Betamethasone valerate lotion 0.1% BD Placebo lotion BD Findings reported for foam and lotion combined: Betamethasone (B) Placebo (P) 	
Outcomes	 Erythema, scaling, thickness, pruritis IAGI (7-pt: worse to completely clear) PAGI (7-pt: worse to completely clear) 	
Notes	Connectics Corporation sponsored by the trial. This was a scalp trial. There was SD imputation (TSS).	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Franz 1999 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	9.5%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Franz 2000

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 188 Treatment duration: 2 wks; FU: 4 wks LF: 0 (0%) BC: unclear Age: not reported Gender (per cent men): 49.5% Severity: mean TSS (0 to 12) = 7.25 INCLUSION CRITERIA • Moderate to severe scalp psoriasis (each of 3 primary signs ≥ 2) • Scalp involvement $\geq 10\%$ • Adults EXCLUSION CRITERIA • Not reported
Interventions	 Clobetasol propionate foam 0.05% BD Placebo foam BD Clobetasol propionate lotion 0.05% BD Placebo lotion BD Findings reported for foam and lotion combined: Clobetasol (C) Placebo (P)

Franz 2000 (Continued)

Outcomes	 Signs: erythema, scaling, thickness, pruritis TSS (0 to 12) IAGI (7-pt: worse to completely clear) PAGI (7-pt: worse to completely clear)
Notes	Connectics Corporation sponsored by the trial. This was a scalp trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Geilen 2000

Methods	DESIGN Within-patient Nurse delivery ALLOCATION Random
	Method of randomisation: not reported Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 7 Treatment duration: 3 wks; FU: 3 wks LF: 0 (0%) BC: yes Age: 36 to 67 Gender (per cent men): 100% Severity: TSS (0 to 8) = 6.72 (0.76SD)

	INCLUSION CRITERIA • Chronic plaque psoriasis EXCLUSION CRITERIA • Not reported
Interventions	 Mycophenolic acid ointment 1% plus occlusion OD (M) Placebo ointment plus occlusion OD (P)
Outcomes	1. TSS (erythema induration) (0 to 8)
Notes	The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Gottlieb 2003

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Conducted in 17 centres; stratification not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 279
- un cherpenne	Treatment duration: 2 wks; FU: 4 wks
	LF: 8 (2.9%)

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.

Gottlieb 2003 (Continued)

Loss to follow up	Low risk	-
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	This was partially demonstrated.

Grattan 1997 (H)

Methods	DESIGN Within-patient Delivery unclear ALLOCATION Random Method of randomisation: pre-determined randomisation schedule Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Described
Participants	 N: 25 Treatment duration: 4 wks; FU: 16 weeks LF: not reported BC: yes Age: 44.0 (range = 20 to 72) Gender (per cent men): 52.0%) Severity: BSA = 16.1% (range = 4.1% to 47.8%); TSS (target sites) = 6.3 (range = NR) INCLUSION CRITERIA Bilateral stable chronic plaque psoriasis Adult Hospitalised for routine dithranol treatment EXCLUSION CRITERIA Intolerance of dithranol Unstable or pustular psoriasis Calcium metabolism disorders Systemic psoriasis treatment Recent UVB or PUVA therapy Pregnancy or lactation
Interventions	 Calcipotriol ointment 0.005% BD (C) Dithranol in aqueous gel (dose titration 0.1% to 2.0%) BD (D)
Outcomes	 Severity (erythema; scaling; palpability) Total Severity Score Patient assessment of irritation (VAS) Investigator assessment of skin staining (none, mild, moderate, or severe)

Grattan 1997 (H) (Continued)

Notes	There was inpatient treatment to ensure high level of compliance. Dermal Laboratories sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Grattan 1997 (P)

Methods	DESIGN
	Within-patient
	Delivery unclear
	ALLOCATION
	Random
	Method of randomisation: pre-determined randomisation schedule
	Concealment: unclear
	BLINDING
	Open
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 12 Treatment duration: 4 wks: FU: 16 wks LF: 0 (0%) BC: yes Age: 50.3 (range = 33 to 75) Gender (per cent men): 33% (4/12) Severity: BSA 17.1% (range = 4.7% to 45.7%); TSS = 6.3 (range = 5 to 7) INCLUSION CRITERIA • Bilateral stable chronic plaque psoriasis • Adult • Hospitalised for routine dithranol treatment EXCLUSION CRITERIA

Grattan 1997 (P) (Continued)

	 Intolerance of dithranol Unstable or pustular psoriasis Calcium metabolism disorders Systemic psoriasis treatment Recent UVB or PUVA therapy Pregnancy or lactation
Interventions	 Dithranol in aqueous gel (dose titration 0.1 to 2.0%) BD (D) Placebo (vehicle) (P)
Outcomes	 Severity (erythema; scaling; palpability) Total Severity Score, patient assessment of irritation (VAS) Investigator assessment of skin staining (none, mild, moderate, or severe)
Notes	There was inpatient treatment to ensure high level of compliance. Dermal Laboratories sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Green 1994

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 49 Treatment duration: 4 wks; FU: 4 wks LF: 3 (6.1%) BC: unclear Age: not reported Gender (per cent men): not reported Severity: mean TSS (0 to 12) = 6.7 INCLUSION CRITERIA Mild to moderate scalp psoriasis and a history of psoriasis elsewhere on the body Adult EXCLUSION CRITERIA Excessively thick scalp psoriasis Other scalp disease Marked deterioration of scalp psoriasis at entry Recent systemic or UV therapy Concurrent topical corticosteroid use Vitamin D or calcium supplement Medications that could affect the course of the disease Hypercalcaemia Hepatic or renal disease At risk of pregnancy
Interventions	 Calcipotriol solution 50 mcg/ml BD (C) Placebo (vehicle) (P)
Outcomes	 Signs (erythema; thickness; scaliness; flaking; itching) Total Sign Score (redness, thickness, scaliness) Investigator Global Assessment Patient Global Assessment
Notes	Leo Pharmaceutical Products sponsored the trial. Compliance was assessed by unused medication returned at each visit. The compliance rate for participants in each group was > 90%. This was a scalp trial.

Risk of bias

Green 1994 (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	6.1%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Greenspan 1993

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 80
Tarticipants	Treatment duration: 3 wks; FU: 3 wks
	LF: 9 (11.3%)
	BC: unclear
	Age: 51.5 (range = 20 to 77)
	Gender (per cent men): 43.8%
	Severity: not reported
	INCLUSION CRITERIA
	• Mild to moderate psoriasis
	EXCLUSION CRITERIA
	• Recent systemic or topical treatment for psoriasis
	Contraindication to low-potency corticosteroids
	Pregnant, nursing, or planning pregnancy
Interventions	• Desonide lotion 0.05% TDS (DL)
	• Desonide cream 0.05% TDS (DC)
	• Placebo (vehicle lotion) (P)

Greenspan 1993 (Continued)

Outcomes	 Severity (erythema; scaling; induration; pruritis) Investigator Global Assessment
Notes	The trial did not report sponsorship, but 3 of the authors were employed by Owen/ Galderma laboratories. There was SD imputation (IAGI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	11.3%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Gribetz 2004

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: 'validated system that automates the random assignment of treatment codes' Concealment: adequate BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 57 Treatment duration: 8 wks; FU: 8 wks LF: 6 (10.5%) BC: yes Age: 47.8 (range = 21 to 88) Gender (per cent men): 50.9% Severity: PGA, per cent moderate = 72%; PGA, per cent severe = 29.8%; TSS = 5.34 (range = 3.0 to 9.0)

	 INCLUSION CRITERIA Stable chronic plaque psoriasis Moderate to severe inverse psoriasis affecting axillae, inguinal, inframammary, or gluteal cleft regions (duration ≥ 6 mths) PGA ≥ 3 Erythema ≥ 2 Aged ≥ 18 EXCLUSION CRITERIA Clinically significant laboratory abnormalities Hypersensitivity to study drug or vehicle Systemic, phototherapy, or immuno-modifying agents within previous 30 dys Topical therapies within previous 14 dys Unstable plaque psoriasis, pustular, drug associated or erythrodermic psoriasis
Interventions	 Pimecrolimus cream (Elidel®) 1% BD (PM) Placebo cream BD (P)
Outcomes	 Investigator's Global Assessment of overall severity (PGA) (5-pt: clear to severe disease) Target Area Score (TSS) (erythema, induration, scaling) (0 to 9) Patient Self-Assessment (control of psoriasis over previous 1 wk) (4-pt: 0 = complete control; 3 = uncontrolled)
Notes	Novartis Pharmaceuticals Group sponsored the trial. No instances of skin atrophy were reported. This was an inverse psoriasis trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	Novartis supplied identical tubes identi- fied only by randomisation number (par- ticipants and investigators blinded to tube contents)
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was by a validated system that automated the random assignment of treatment codes
Loss to follow up	Low risk	10.5%
Baseline assessments	Low risk	These were reported.

Gribetz 2004 (Continued)

Baseline comparability demonstrated	Low risk -
Guenther 2000	
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 120 Treatment duration: 8 wks; FU: 20 wks LF: 14 (11.7%) BC: yes Age: 48.5 Gender (per cent men): 60.8% Severity: not reported INCLUSION CRITERIA Stable chronic plaque psoriasis BSA involvement between 5% and 20% Adult EXCLUSION CRITERIA Pregnancy Lactation Unreliable contraception Unstable plaque psoriasis Other types of psoriasis or other concomitant dermatological disorder Hypercalaemia Uncontrolled systemic disease Likelihood of prolonged UV exposure Concomitant systemic or topical therapies that might affect psoriasis Adherence to washout requirements
Interventions	 Tazarotene gel, 0.1% ON, plus mometasone furoate cream 0.1% OM (TM) Calcipotriol ointment 0.005% BD (C) 12 weeks' maintenance for both groups with emollient only.
Outcomes	 IAGI (0 to 6; exacerbation to complete clearance) Erythema, scaling, thickness (0 to 4 for each) BSA involvement Patient assessments (efficacy, comfort of skin; outlook for long-term control; overall impression of treatment)

Guenther 2000 (Continued)

	5. Adverse events	
Notes	Allergan Inc. sponsored the trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	11.7%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Guenther 2002 (H)

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated random numbers table
	Concealment: adequate
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 828
Tarticipants	Treatment duration: 4 wks; FU: 4 wks
	LF: 10 (1.2%)
	BC: yes
	Age: 48.5 (14.3SD)
	Gender (per cent men): 64.0%
	Severity: mean PASI = 10.5; mean duration psoriasis = 18.3 yrs
	INCLUSION CRITERIA
	Aged 18 to 86
	Chronic plaque psoriasis
	• BSA involvement $\geq 10\%$
	EXCLUSION CRITERIA

Guenther 2002 (H) (Continued)

	 Systemic therapy within previous 6 wks Topical antipsoriatic therapy within previous 2 wks Concurrent use of type III/IV topical corticosteroids Recent UV exposure Pregnancy Lactation Concurrent use of other medicines that could affect course of psoriasis
Interventions	 Calcipotriol (50 mcg/g) and betamethasone dipropionate (0.5 mg/g) ointment ON, plus placebo OM (D1) Calcipotriol (50 mcg/g) and betamethasone dipropionate (0.5 mg/g) ointment BD (D2) Calcipotriol BD (C) Placebo BD (P)
Outcomes	 PASI (head excluded) IAGI (6-pt: worse to clearance) PAGI (6-pt: worse to clearance) Percentage change in thickness score Speed of response (PASI) at 1 week Adverse events Quality of life: Psoriasis Disability Index; EQ-5D and EQ-VAS (reported in van de Kerkhof 2004)
Notes	Leo Pharmaceuticals sponsored the trial. The trial was conducted across 57 centres in 8 countries.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	All study personnel and participants were blinded to treatment assignment for the du- ration of the study
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	1.2%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Guenther 2002 (P)

Methods	DESIGN Between-patient
	Participant delivery ALLOCATION
	Random
	Method of randomisation: computer-generated random numbers table Concealment: adequate
	BLINDING Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 828 Treatment duration: 4 wks; FU: 4 wks
	LF: 10 (1.2%)
	BC: yes
	Age: 48.5 (14.3SD)
	Gender (per cent men): 64.0% Severity: mean PASI = 10.5; mean duration psoriasis = 18.3 yrs
	INCLUSION CRITERIA
	• Aged 18 to 86
	Chronic plaque psoriasis
	• BSA involvement $\geq 10\%$ EXCLUSION CRITERIA
	Systemic therapy within previous 6 wks
	• Topical antipsoriatic therapy within previous 2 wks
	Concurrent use of type III/IV topical corticosteroids
	 Recent UV exposure Pregnancy
	Lactation
	• Concurrent use of other medicines that could affect course of psoriasis
Interventions	• Calcipotriol (50 mcg/g) and betamethasone diproprionate (0.5 mg/g) ointment
	ON, plus placebo, OM (D1)
	• Calcipotriol (50 mcg/g) and betamethasone diproprionate (0.5 mg/g) ointment BD (D2)
	Calcipotriol BD (C)
	Placebo BD (P)
Outcomes	1. PASI (head excluded)
	2. IAGI (6-pt: worse to clearance)
	3. PAGI (6-pt: worse to clearance)
	 Percentage change in thickness score Speed of response (PASI) at 1 week
	6. Adverse events
	7. Quality of life: Psoriasis Disability Index; EQ-5D and EQ-VAS (reported in van de Kerkhof 2004)
Notes	Leo Pharmaceuticals sponsored the trial.
	The trial was conducted across 57 centres in 8 countries.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	All study personnel and participants were blinded to treatment assignment for the du- ration of the study
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	1.2%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Han 2001

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Open
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 208
T articipants	Treatment duration: 12 wks; FU: 12 wks
	LF: 9 (4.3%)
	BC: yes
	Age: 40.4 (8.9SD)
	Gender (per cent men): 59.8%
	Severity: TSS (0 to 20) = $16.1 (11.0$ SD)
	Duration (yrs): 9.4 (8.9SD) INCLUSION CRITERIA
	People with chronic plaque psoriasis
	 Aged between 18 and 65 BSA between 2% and 30%
	• BSA between 2% and 50% EXCLUSION CRITERIA
	Known allergy to study drug constituents

Han 2001 (Continued)

	 History of other skin diseases Pustular or erythrodermic psoriasis Topical treatments within previous 2 wks PUVA within previous 4 wks UVB within previous 2 wks Alcohol or drug abuse Renal, hepatic, or immunity disorder Pregnancy or risk thereof Lactation Participation in another clinical trial within previous 4 wks Other morbidity likely to affect outcome or raise safety issues
Interventions	 Tazarotene gel 0.05% OD (T) Calcipotriol ointment 5 mcg/g BD (C)
Outcomes	 TSS (0 to 20) Proportion of patients achieving effective response Curative rate
Notes	The trial did not report sponsorship. Translation support was received for data extraction.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	4.3%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Harrington 1996a

Methods	DESIGN	
withous	Between-patient	
	Participant delivery	
	ALLOCATION	
	Random	
	Method of randomisation: not reported	
	Concealment: unclear	
	BLINDING	
	Double-blind (participant/investigator)	
	WITHDRAWAL/DROPOUT	
	Described	
Participants	N: 413	
-	Treatment duration: 8 wks; FU: 8 wks	
	LF: 47 (11.4%)	
	BC: yes, except average age in placebo grou	Ip higher than for A and B ($P = 0.02$)
	Age: 44.6	
	Gender (per cent men): 52.8% Severity: PASI (modified) = 8.3 (range = 0.6 to 59.4) Duration (yrs): 17.7 (range = 0.04 to 70) INCLUSION CRITERIA • Stable chronic plaque psoriasis on trunk or limbs • Adult	
	EXCLUSION CRITERIA	
	Recent systemic medication or photon	herapy for psoriasis
	 Hepatic or renal disease Raised serum calcium Calcium supplements or vitamin D 	
Interventions	• Calcipotriol cream 50 mcg/g BD as follows:	
incrventions	 Catciportion cream 50 integrg DD as it Cream A (dissolved) (CA) 	JIOWS.
	• Cream B (suspended) (CB)	
	 Placebo (vehicle of A) (P) 	
Outcomes	1. PASI (modified to exclude head)	
	 Investigator Global Assessment Patient Global Assessment 	
	5. Patient Global Assessment	
Notes	Leo Pharmaceuticals sponsored the trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

 Blinding (performance bias and detection bias)
 Low risk
 The trial was double-blind (participant/investigator).

 All outcomes
 House
 House

Topical treatments for chronic plaque psoriasis (Review)

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Harrington 1996a (Continued)

Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	11.4%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Helfrich 2007

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: unclear (method inadequately described) Concealment: unclear (inadequately described) BLINDING Double-blind (participant/investigator/assessor) WITHDRAWAL/DROPOUT Described
Participants	N: 185 Treatment duration: 8 wks; FU: 8 wks LF: 3 (1.6%) BC: yes Age: 49.0 (range = 19 to 83) Gender (per cent men): 56.8% Severity: PGA = 3.2 (range = 3 to 4); PSS = 8.1 (range = 4.7 to 12) Ethnicity (per cent white): NR Duration: 17.3 (range = 0.5 to 65) INCLUSION CRITERIA • People aged 18 and over with chronic plaque psoriasis affecting 2% to 10% BSA with severity "appropriate" for topical therapy • PGA score: > = 3 EXCLUSION CRITERIA • Topical therapies within previous 2 wks • Systemic/UV therapy within previous 4 wks • Biological within previous 6 mths • Oral vitamin D (> 400 IU/day), oral calcium (> 1200 mg/day) within previous 30 days • Pregnancy or risk thereof • Lactation • Significant medial comorbidity • Sensitivity to study drug • Concomitant antipsoriatics therapy, immunosuppressive drugs, lithium, hydoxycholoroquine, or biologicals • Non-compliance with dosing of study medication

Helfrich 2007 (Continued)

Interventions	 Becocalcidiol ointment 75 mcg/g BD (B2) Becocalcidiol ointment 75 mcg/g ON and placebo OM (B1) Placebo ointment BD (P) Max: 8 g ointment/day Participants were permitted concurrent oral vitamin D (= < 400 IU/day), oral calcium (< = 1200 mg/day), or both; or tar shampoo for scalp psoriasis
Outcomes	 Physician's Static Global Assessment of Overall Lesion Severity (PGA) (6-point scale: clear to very severe) Psoriasis Symptom Severity (PSS) score (0 to 12): sum of scores for erythema, scaling, induration (each scored 0 to 4) Adverse events Laboratory evaluations Compliance (participants who missed > 6 consecutive doses discontinued the study)
Notes	QuatRx Pharmaceuticals (product now owned by Deltanoid Pharmaceuticals) sponsored the trial The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant and investigator/assessor)
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	1.6%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated (clinical and demo- graphic).

Henneicke-v. Z. 1993

Methods	DESIGN Within-patient (placebo) Between-patient (active) Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 73 Treatment duration: 8 wks; FU: 8 wks LF: 21 (28.8%) BC: yes Age: 40.5 (median); range = 18 to 71 (N = 52) Gender (per cent men): 65.8% (N = 73) Severity: TSS (0 to 12) = 7.4 INCLUSION CRITERIA • Aged 18 to 71 • Moderate plaque psoriasis • $9 > TSS \ge 4$ EXCLUSION CRITERIA • Systemic antipsoriatic drugs within previous 4 wks • UV therapy within previous wk • Drug-induced psoriasis • Cancer • Pregnancy • Lactation • Severe organ dysfunction • Metabolic disorders • Abuse of drugs or alcohol
Interventions	 Omega-3-polyunsaturated fatty acids ointment 1% BD (O3(1)) Placebo (vehicle) 1% BD Omega-3-polyunsaturated fatty acids ointment 10% BD (O3(10)) Placebo (vehicle) 10% BD (P)
Outcomes	 Local psoriasis severity index (TSS equivalent): erythema, induration, desquamation Area of indicator lesion Pruritis Investigator's subjective intra-individual comparison (left side better than right side) Patient's subjective intra-individual comparison (left side better than right side) Compliance: tube weight compared with expected use

Henneicke-v. Z. 1993 (Continued)

Notes	The trial did not report sponsorship.	
Risk of bias		
Bias	Authors' judgement Support for judgement	
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	High risk	28.8%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Highton 1995

Methods	DESIGN	
	Between-patient	
	Participant delivery	
	ALLOCATION	
	Random	
	Method of randomisation: not reported	
	Concealment: unclear	
	BLINDING	
	Double-blind (participant/investigator)	
	WITHDRAWAL/DROPOUT	
	Described	
Participants	N: 277	
Participants	Treatment duration: 8 wks	
	LF: 30 (10.8%)	
	BC: clinical severity comparable, demographics unclear	
	Age: not reported	
	Gender (per cent men): not reported	
	Severity: TSS (0 to 8) = 3.90 ; BSA = 9.1%	
	INCLUSION CRITERIA	
	Moderately severe stable plaque psoriasis	
	• Plaque elevation score greater than or equal to 4 (0 to 8)	
	 Not pregnant or nursing during the duration of the study EXCLUSION CRITERIA 	
	• Recent topical or systemic psoriasis treatment, prolonged exposure to sunlight,	

Highton 1995 (Continued)

	 phototherapy Photochemotherapy Hypercalcemia Erythrodermic or pustular psoriasis Calcium, vitamin A or D supplements 	
Interventions	 Calcipotriene ointment 0.005% BD (C) Placebo (vehicle) (P) 	
Outcomes	 Severity (erythema; plaque elevation; scaling; overall disease severity) 75% improvement scores Investigator Global Assessment (7-pt: worse to completely clear) 	
Notes	Bristol Myers Squibb sponsored the trial. There was SD imputation (TSS).	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection	Low risk	The trial was double-blind (participant/in-

bias) All outcomes		vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	10.8%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Hindsén 2006 (H)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Not described
Participants	N: 360 Treatment duration: 2 or 6 wks; FU: 6 or 10 wks LF: NR BC: not demonstrated Age: NR Gender (per cent men): NR Severity: NR INCLUSION CRITERIA • People with moderately severe plaque psoriasis on elbows, knees, or both EXCLUSION CRITERIA • Not stated
Interventions	 Calcipotriol ointment 50 mcg/g BD (no occlusion) for 6 wks (C) Calcipotriol ointment 50 mcg/g plus occlusion, once/week for 2 wks (CO) [Placebo ointment plus occlusion, once/week for 2 wks (P)]
Outcomes	1. Total Sign Score (redness, thickness, scaliness) (TSS): 0 to 24
Notes	Leo Pharma, Ballerup, sponsored the trial. It was unclear how participants were blinded to treatment options where 2/3 involved occlusion The sponsor supplied unpublished data.

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial was reported to be double-blind (participant/investigator), but it was un- clear how they blinded participants to treat- ment options where 2/3 involved occlusion
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.

Hindsén 2006 (H) (Continued)

Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	This was not demonstrated.

Hindsén 2006 (P)

Bias	Authors' judgement	Support for judgement	
Risk of bias			
Notes	It was unclear how participants occlusion	Leo Pharma, Ballerup, sponsored the trial. It was unclear how participants were blinded to treatment options where 2/3 involved occlusion The sponsor supplied unpublished data.	
Outcomes	1. Total Sign Score (redness,	1. Total Sign Score (redness, thickness, scaliness) (TSS): 0 to 24	
Interventions	Calcipotriol ointment 50	 Calcipotriol ointment 50 mcg/g, BD (no occlusion) for 6 wks (C) Calcipotriol ointment 50 mcg/g plus occlusion, once/week for 2 wks (CO) Placebo ointment plus occlusion, once/week for 2 wks (P) 	
Participants	LF: NR BC: not demonstrated Age: NR Gender (per cent men): NR Severity: NR INCLUSION CRITERIA	Treatment duration: 2 or 6 wks; FU: 6 or 10 wks LF: NR BC: not demonstrated Age: NR Gender (per cent men): NR Severity: NR INCLUSION CRITERIA • People with moderately severe plaque psoriasis on elbows, knees, or both EXCLUSION CRITERIA	
Methods	Between-patient Participant delivery ALLOCATION Random Method of randomisation: not s Concealment: unclear BLINDING	Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT	

Dias	Authors Judgement	Support for Judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Hindsén 2006 (P) (Continued)

Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial was reported to be double-blind (participant/investigator), but it was un- clear how they blinded participants to treat- ment options where 2/3 involved occlusion
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	This was not demonstrated.

Holick 2003

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 15 Treatment duration: 8 wks; FU: 8 wks LF: 0 (0%) BC: not reported Age: 56 (range = 25 to 74) Gender (per cent men): 67 % Severity: not reported INCLUSION CRITERIA Chronic plaque psoriasis 2 symmetrically comparable plaques Failure to respond to at least 1 standard treatment EXCLUSION CRITERIA Kidney disease, hypercalcaemia, hypercalciuria Systemic therapy within previous 30 days Topical therapy within previous 14 days Concomitant medications that interfere with calcium metabolism
Interventions	 Parathyroid hormone (PTH) (1 to 34) 20 mcg/g in Novasome A® liposomal cream BD (PTH) Novasome A® liposomal cream BD (P)

Holick 2003 (Continued)

Outcomes	 Global severity score (0 to 24) PASI (for open trial phase only)
Notes	The National Institutes for Health (Department of Health and Human Services) and the Department of Defense Small Business Innovation Research (SBIR) Program sponsored the trial through grants. The trial also reported a uncontrolled open, large area, study for N = 10 (PTH applied OD for up to 11 months)

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Huang 2009

Methods	DESIGN	
	Between-patient	
	Participant delivery	
	ALLOCATION	
	Random	
	Method of randomisation: unclear	
	Concealment: unclear	
	BLINDING	
	Double-blind (participant/investigator)	
	WITHDRAWAL/DROPOUT	
	Described	
Participants	N: 320	
rarticipants	Treatment duration: 4 wks; FU: 4 wks	
	LF: 0 (0%)	
	BC: stated no significant difference in clinical or demographic characteristics (data not	
	shown)	
	Age: NR	
	150.111	

	Gender (per cent men): NR
	Severity: NR
	INCLUSION CRITERIA
	• Chinese people aged 18 to 65 with stable plaque psoriasis
	• At least 1 area (or arm, leg, body) with mPASI score > = 2 (i.e. % BSA > = 10% of
	affected area)
	EXCLUSION CRITERIA
	Psoriasis of the face
	• Severe disease requiring steroids or other systemic medication
	• % BSA > = 30%
	• Use of systemics during previous 4 weeks, topicals during previous 2 weeks
	Participation in another trial during previous 3 months
	• Iodine deficiency, hypercalcaemia, Cushing's Syndrome, diabetes, or other
	metabolic disease
	• Heart disease, kidney disease, liver disease
	• Immunesuppression
	• Cancer
	Mental disorder
	• Failure to consent
	Pregnancy or risk thereof
	Lactation
	Investigator opinion of unsuitability
Interventions	• Calcipotriol ointment BD (C)
	• Calcipotriol betamethasone ointment ON, placebo OM (C-B)
	Participant gave written consent to use < = 100 g/wk.
Outcomes	1. mPASI (0 to 64.8)
	2. Treatment success: PASI75
	3. TSS (redness, thickness, scaliness) (0 to 12)
	4. DLQI
	5. Adverse events
	6. Laboratory tests: urine, blood, biochemistry, liver function, electrolytes, serum
	calcium
	7. Atrophy, telangiectasias
Notes	The trial did not report sponsorship.
	We received translation support from native speaker.
	· · · · · · · · · · · · · · · · · · ·

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Huang 2009 (Continued)

Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Unclear risk	The trial authors state these were under- taken.
Baseline comparability demonstrated	Unclear risk	The trial authors state that groups were comparable at baseline (not demonstrated)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Described
Participants	 N: 114 Treatment duration: 8 wks; FU: 8 wks LF: 28 (24.6%) BC: yes Age: 42.3 Gender (per cent men): 74.4% Severity: PASI (mean) = 11.8 Duration of psoriasis (mths) (mean): 185.1 (range = 1 to 85) INCLUSION CRITERIA Chronic plaque psoriasis of at least moderate severity Aged over 18 White or Asian origin EXCLUSION CRITERIA Systemic or intralesional therapy or photo-chemotherapy within previous 2 mths Topical antipsoriatics within previous wk or concomitant Other medications that could affect psoriasis Pregnancy Inadequate contraception
Interventions	 Calcitriol ointment 3 mcg/g BD (C) Short contact dithranol 0.25 to 2% OD (D)

Hutchinson 2000 (Continued)

Outcomes	 PASI IAGI (6-pt: worse to clearing) Overall global severity (5-pt: none to very severe) Psoriasis Disability Index (quality of life) (scale NR) Cosmetic acceptability (1 = good/none to 3 = not acceptable): staining, irritation Adverse events
Notes	The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	24.6%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Jarratt 2004

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated list
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Unclear
Danticipanto	N: 142
Participants	Treatment duration: 4 wks; FU: 6 wks
	LF: 1 (0.7%)
	BC: yes
	Age: 45.1 (15.37SD)
	Gender (per cent men): 42.3%
	Severity: TSS (0 to 9) = 6.6 ; GSS (6-pt (rescaled: 0 = very severe to 5 = clear) = 1.65 (0.

	61SD), N = 142
	INCLUSION CRITERIA
	• Aged 12 or over
	• Moderate to severe scalp psoriasis (global severity score ≥ 3)
	• Compliance with wash-out periods for systemic therapies (details not reported)
	EXCLUSION CRITERIA
	Pregnancy or risk thereof
	 Known allergy to test products
	 Need for systemic therapy or other concomitant antipsoriatics
	• Excessive UV exposure
Interventions	 Clobetasol propionate shampoo 0.05% OLUX® OD (C) Placebo shampoo OD (P) Treatments applied OD, left to dry for 15 minutes, then washed out
	Placebo represented by vehicle for clobetasol propionate
Outcomes	 Global severity score (GSS) (6-pt: 0 = clear to 5 = very severe) Total Severity Score (erythema, thickening, scaling) (TSS) (0 to 9) Individual sign scores for erythema, thickening, scaling pruritis, per cent scalp involvement (4-pt: 0 = none to 3 = severe) IAGI (5-pt: worse to clear) PAGI (5-pt: worse to clear)
Notes	Galderma R&D Inc sponsored the trial. Missing data imputed using last observation carried forward. No cases of skin atrophy, teleangiectasia, or acne were reported. This was a scalp trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	-
Loss to follow up	Low risk	0.7%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Jarratt 2006

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 120 Treatment duration: 4 wks; FU: 8 wks LF: 0 (0%) BC: yes Age: 48.0 (12.9SD) Gender (per cent men): 60.0% Severity: % BSA = 7.69% (6.17%SD); ODS (moderate) = 90.8%; ODS (severe/very severe) = 9.2% Ethnicity (per cent white): 94.2% Duration: NR INCLUSION CRITERIA People aged 18 and over with moderate to severe chronic plaque psoriasis BSA > = 2% (excluding face, scalp, groin, axillae, and other intertriginous areas) Overall disease severity score (0 to 4) > = 3 EXCLUSION CRITERIA Non-compliance with treatment specific wash-out periods on topical and systemic treatments or phototherapy or natural UV light (periods NS) Pregnancy or risk thereof
Interventions	 Clobetasol propionate 0.05% spray BD (CP) Placebo spray BD (P) Then 4-week treatment-free follow-up period. There was 8 hrs between applications, and the treated area was unwashed for 4 hours after application
Outcomes	 Scaling, erythema, plaque elevation, pruritus, overall disease severity (ODS) each scored on 5-pt scale (0 = clear to 4 = severe/very severe) Success rate: ODS < = 2 at wk 2 and ODS < = 1 at wk 4 ODS at wk 1 and wk 8 Signs at wk 1 and wk 8 ODS < = 1 at wk 8 Adverse events (atrophy, telangiectasia, burning/stinging, folliculitis); HPA axis suppression
Notes	Galderma Pharmaceuticals sponsored the trial. The trial also reported CIC data on a safety RCT (2 or 4 wks with clobetasol propionate 0.05% spray, < = 50 g/wk): 8/36 had HPA suppression; all returned to normal within 2 wks of end of treatment (EOT)

Jarratt 2006 (Continued)

The trialist supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	The trial reported these.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Jekler 1992

Methods	DESIGN
Wethous	Within-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 30
1	Treatment duration: 8 wks
	LF: 3 (10%)
	BC: not reported
	Age: 45.2 (14.0SD; N = 27)
	Gender (per cent men): 77.8%; N = 27
	Severity: severity (mean score, 0 to 3) = 1.9 ; N = 27
	Duration disease (years): 20.5 (14.8SD; $N = 27$)
	Duration exacerbation (mths): $5.1 (6.4SD; N = 27)$
	INCLUSION CRITERIA
	Chronic plaque-type psoriasis with bilateral lesions of equal clinical severity
	 Adult.
	EXCLUSION CRITERIA
	Topical or systemic corticosteroids

Jekler 1992 (Continued)

	• Recent phototherapy	
Interventions	Dithranol 2% ointment 1 minute thePlacebo (vehicle) (P)	rapy OD (D)
Outcomes	 Severity (pruritis; erythema; scaling; in Investigator's assessment of degree of c Patient's assessment of degree of clearing 	clearing
Notes	E Merck AB sponsored the trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

All outcomes		
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	10.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Jemec 2008 (H)

Methods	DESIGN
methods	Between-patient
	•
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: pre-planned computer-generated randomisation code list (2:
	4:4:1)
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
	N 1505
Participants	N: 1505
	Treatment duration: 8 wks; FU: 8 wks
	LF: 32 (2.1%)
	BC: yes
	<i>b</i> o. <i>y</i> o

	Age: 49.0 (15.8SD); range = 17 to 97 Gender (per cent men): 44.8% Ethnicity (per cent white): 96.3% Severity: IGA mild disease = 6.5%; IGA moderate disease = 56.2%; IGA severe disease = 31.6%; IGA very severe disease = 5.7%; TSS (0 to 12) = 6.82 (1.85SD) Duration (yrs): 16.5 (13.6SD) INCLUSION CRITERIA • People aged > = 18 with scalp plaque psoriasis involving > = 10% scalp • Amenable to topical treatment with < = 100 g medication/wk • Diagnosis of psoriasis vulgaris on trunk/limbs • Score at least moderate in 1 sign (erythema, thickness, scaliness) and at least slight for other signs • IGA mild to very severe EXCLUSION CRITERIA • PUVA or grenz ray or relevant systemic therapy within previous 4 wks • UVB or topical scalp antipsoriatic therapy or topical very potent (WHO group IV) corticosteroid on face/body within previous 2 wks • Biological therapy within previous 6 mths • Planned initiation of/changes to concomitant medication that could affect scalp psoriasis • Planned exposure to sun • Erythrodermic, exfoliative, or postural psoriasis • Viral lesions • Skin infections • Atrophic skin on scalp • Known/suspected abnormality of calcium homeostasis • Severe renal insufficiency
Interventions	 Severe hepatic disorder Calcipotriol 50 mcg/g gel OD (C) Betamethasone dipropionate 0.5 mg/g gel OD (B) Combined gel: calcipotriol 50 mcg/g + betamethasone dipropionate 0.5 mg/g OD (C-B) Placebo gel OD (P)
Outcomes	 Total Sign Score (0 to 12) Investigator's Global Assessment of Disease Severity of the scalp (IGA): 6-pt: absence of disease to very severe disease Treatment success: IGA absence of disease or very mild disease Patient overall assessment of treatment response on a 7-pt scale (worse, unchanged, slight improvement, moderate improvement, marked improvement, almost clear, cleared) Compliance (self-report) Adverse events Laboratory tests (serum calcium, serum albumin)
Notes	Leo Pharma A/S, Ballerup, Denmark, sponsored the trial. The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	A preplanned computer-generated ran- domisation code list was used
Loss to follow up	Low risk	2.1%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Jemec 2008 (P)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: pre-planned computer-generated randomisation code list (2: 4:4:1) Concealment: unclear BLINDING Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT Described
Participants	N: 1505 Treatment duration: 8 wks; FU: 8 wks LF: 32 (2.1%) BC: yes Age: 49.0 (15.8SD); range = 17 to 97 Gender (per cent men): 44.8% Ethnicity (% white): 96.3% Severity: IGA mild disease = 6.5%; IGA moderate disease = 56.2%; IGA severe disease = 31.6%; IGA very severe disease = 5.7%; TSS (0 to 12) = 6.82 (1.85SD) Duration (yrs): 16.5 (13.6SD) INCLUSION CRITERIA • People aged > = 18 with scalp plaque psoriasis involving > = 10% scalp • Amenable to topical treatment with < = 100 g medication/wk • Diagnosis of psoriasis vulgaris on trunk/limbs

	 Score at least moderate in 1 sign (erythema, thickness, scaliness) and at least slight for other signs IGA mild to very severe EXCLUSION CRITERIA PUVA or grenz ray or relevant systemic therapy within previous 4 wks UVB or topical scalp antipsoriatic therapy or topical very potent (WHO group IV) corticosteroid on face/body within previous 2 wks Biological therapy within previous 6 mths Planned initiation of/changes to concomitant medication that could affect scalp psoriasis Planned exposure to sun Erythrodermic, exfoliative, or postural psoriasis Viral lesions Skin infections Atrophic skin on scalp Known/suspected abnormality of calcium homeostasis Severe renal insufficiency Severe hepatic disorder
Interventions	 Calcipotriol 50 mcg/g gel OD (C) Betamethasone dipropionate 0.5 mg/g gel OD (B) Combined gel: calcipotriol 50 mcg/g + betamethasone dipropionate 0.5 mg/g OD (C-B) Placebo gel OD (P)
Outcomes	 Total Sign Score (0 to 12) Investigator's Global Assessment of Disease Severity of the scalp (IGA): 6-pt: absence of disease to very severe disease Treatment success: IGA absence of disease or very mild disease Patient overall assessment of treatment response on a 7-pt scale (worse, unchanged, slight improvement, moderate improvement, marked improvement, almost clear, cleared) Compliance (self-report) Adverse events Laboratory tests (serum calcium, serum albumin)
Notes	Leo Pharma A/S, Ballerup, Denmark, sponsored the trial. The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Jemec 2008 (P) (Continued)

Randomisation method reported	Low risk	A preplanned computer-generated ran- domisation code list was used
Loss to follow up	Low risk	2.1%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Ji 2008

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 250 Treatment duration: 12 wks; FU: 12 wks LF: 7 (2.8%) BC: yes Age: 41.7 (12.2SD) Gender (per cent men): 65.6% Ethnicity (per cent Asian): 100% Severity: per cent BSA = 18%, range = 1% to 35%; DSS (0 to 12) = 8.12 (1.97SD), range = 3 to 12 INCLUSION CRITERIA People aged 18 to 65 with stable mild to moderate plaque psoriasis BSA < = 35% Diameter of target lesions > = 3 cm EXCLUSION CRITERIA Pregnancy or risk thereof Acute guttate psoriasis Erythrodermic, pustular, arthropathic psoriasis Hypercalcaemia Renal dysfunction Calcium nephrolithiasis Use of topical antipsoriatic therapy within previous 2 wks Use of systemic antipsoriatic therapy within previous 8 wks

Ji 2008 (Continued)

Interventions	 Calcipotriol 50 mcg/g ointment BD (C1) Calcitriol 3 mcg/g BD (C2) Usage restrictions: < = 210 g/wk calcitriol < = 100 g/wk calcipotriol
Outcomes	 Investigator's Assessment of Global Improvement (IAGI; 4-pt: 0 = worse/no change to 3 = clear/almost clear) Subject's Assessment of Global Improvement (PAGI) (4-pt: 0 = worse/no change to 3 = clear/almost clear) Dermatological Sum Score (DSS): erythema, plaque elevation, scaling: 0 to 12 Local: cutaneous safety (investigator): 5-pt: 0 (none) to 4 (very severe); cutaneous discomfort (subject): 5-pt: 0 (none) to 4 (very severe) Adverse events Systemic: routine blood and urine safety parameters, albumin-adjusted serum total calcium Compliance (medication usage)
Notes	Galderma Laboratories LP sponsored the trial. The trial was set in China. We received translation support for data extraction.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The study was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	2.8%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Jin 2001

Methods	DESIGN	
	Between-patient	
	Participant delivery	
	ALLOCATION	
	Random Mathad of randomization:	
	Method of randomisation: Not stated	
	Concealment: unclear	
	BLINDING	
	Double-blind (participant/investigator)	
	WITHDRAWAL/DROPOUT	
	Described	
Participants	N: 96	
-	Treatment duration: 6 wks; FU: 6 wks	
	LF: 7 (7.3%)	
	BC: yes	
	Age: 35.8 (range = $18 \text{ to } 65$)	
	Gender (per cent men): 57.3% Severity: TSS (0 to 20) = 9.4	
	INCLUSION CRITERIA	
	Chronic plaque psoriasis	
	• Aged over 18? (cannot translate)	
	EXCLUSION CRITERIA	
	• (cannot translate)	
Interventions	Anti-IL-8 monoclonal antibody creanPlacebo (P)	n (M)
0	1. EC	
Outcomes	 Efficacy rate Cure rate 	
	 Cure face Erythema, infiltration, scaling, pruriti 	s
	4. TSS (skin damage, erythema, infiltration, thickness, scaling)	
	5. IAGI (4-pt: failure to cure)	Jan Barris
Notes	Biological Technical Medical Trade Limited	d sponsored the trial
	We received translation support for data ex	-
Risk of bias		
Bias	Authors' judgement	Support for judgement

Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.

Jin 2001 (Continued)

Loss to follow up	Low risk	7.3%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Jorizzo 1997

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 89 Treatment duration: 4 wks; FU: 6 wks LF: Unclear BC: yes Age: 49.7 (range = 21 to 84) Gender (per cent men): 65% Severity: per cent BSA affected = 8.1% Duration of psoriasis (range, years): 1 to 57 Duration of exacerbation (range, wks): 3 to 2080 INCLUSION CRITERIA • Moderate to severe plaque type psoriasis • Non-hospitalised men or non-pregnant; non-lactating women ≥ 12 yrs • Baseline morning serum cortisol concentration of 5 to 18 mcg/100 mL EXCLUSION CRITERIA • Recent topical antipsoriatic medication or other drug that could alter psoriatic status
Interventions	 Clobetasol propionate emollient 0.05% BD (C) Placebo (vehicle) (P)
Outcomes	 Severity (erythema; skin thickening; scaling; pruritis) Total Severity Score (0 to 12) Investigator Global Assessment of improvement (6-pt: worse to cleared) Patient Global Assessment of improvement (5-pt: worse to excellent)
Notes	Glaxo Wellcome Inc. sponsored the trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Jorizzo 2007

Methods	DESIGN
Wethous	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not stated
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Not described
Demisinente	N: 379
Participants	Treatment duration: 6 wks: FU: 6 wks
	LF: NR
	BC: yes (only clinical comparability demonstrated)
	Age: NR
	Gender (per cent men): NR
	Severity: PASI = 7.9 (5.3SD); IGA moderate = 76.2%; IGA severe = 22.2%
	INCLUSION CRITERIA
	People with moderate to severe plaque psoriasis
	EXCLUSION CRITERIA
	Not stated
Interventions	• Calcipotriene 0.005% ointment (C)
	• Calcipotriene 0.005% plus betamethasone dipropionate 0.064% ointment (C-B)
	< = 100 g/wk - the daily dose was not specified.

Jorizzo 2007 (Continued)

Outcomes	 PASI Investigator Global Assessment (IGA) Controlled disease (IGA: absence. very mild disease) Patient Global Assessment of Improvement (PAGI) Treatment success (PAGI marked improvement/cleared)
Notes	Warner-Chilcott sponsored the trial. This was a conference abstract only. We sought data but received no response. There was SD imputation (PASI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Unclear risk	The trial only reported clinical assessments (demographic characteristics not reported)
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated (demo- graphic characteristics not assessed)

Kang 1998

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: identical tubes with computer-generated codes
	Concealment: adequate
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 30
1 articipants	Treatment duration: 6 wks
	incament duration. 0 wks

Kang 1998 (Continued)

	LF: 0 (0%) BC: psoriasis comparable, demographics unclear Age: 41 (range = 18 to 66) Gender (per cent men): 66.7% Severity: TSS (0 to 24) = 12.27 INCLUSION CRITERIA • Mild to moderate stable plaque-type psoriasis • Adult EXCLUSION CRITERIA • Recent systemic therapy, UV, or topical therapy for psoriasis (excluding emollient) • Pregnant or breast-feeding women
Interventions	 Calcipotriene ointment 0.005% BD (C) Placebo (vehicle) (P)
Outcomes	 Signs (erythema; thickness; scaling) TSS (0 to 24) Investigator Global Assessment (7-pt: worse to clear)
Notes	Bristol-Myers Squibb Corporation and from Babcock Dermatologic Endowment (University of Michigan) sponsored the trial through grants

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	Participants were assigned by computer- generated code.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Kanzler 1993

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: identical containers labelled right and left; labelling method not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Not described	
Participants	N: 18 Treatment duration: 4 wks; FU: 4 wks LF: 0 (0%) BC: not reported Age: 45.4 (range = 21 to 66) Gender (per cent men): 55.6% Severity: not reported INCLUSION CRITERIA • Bilaterally similar chronic stable plaque psoriasis EXCLUSION CRITERIA • Recent topical or systemic therapy	
Interventions	 Tar (liquor carbonis detergens) 5% BD (T) Placebo (vehicle) (P) 	
Outcomes	 Severity (erythema; induration; scaling; pruritis) Total Severity Score (0 to 12) Investigator Global Assessment: per cent improvement from baseline, based on TSS 	
Notes	The trial did not report sponsorship. Compliance was assessed by unused medication returned at each visit. The compliance rate for participants in each group was > 90%. This was a scalp trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.

Kanzler 1993 (Continued)

Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Katz 1987a

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 40 Treatment duration: 12 wks; FU: 12 wks LF: 2 (5%) BC: yes Age: 46.7 Gender (per cent men): 60% Severity: per cent participants with BSA affected < 10% = 68% Duration (years): 20.8 INCLUSION CRITERIA • Plaques psoriasis in remission (> 85% resolution) after 2 to 3 weeks treatment with Betamethasone dipropionate Note: 38/59 (64%) achieved remission during the acute phase EXCLUSION CRITERIA • Not achieving remission during acute phase treatment
Interventions	 Betamethasone dipropionate, intermittent maintenance (3 doses at 12-hour intervals each weekend) (B) Placebo (vehicle) (P)
Outcomes	 Signs (erythema; induration; scaling) Area adjusted clinical score Relapse (adjusted clinical score > 35% of baseline score)
Notes	The trial was supported in part by a grant from Schering Plough Corporation. There was SD imputation (TSS).

Risk of bias

Katz 1987a (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	5.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Katz 1991a

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated code
	Concealment unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 94
Tarticipants	Treatment duration: 24 wks; FU: 24 wks
	LF: 4 (4.3%)
	BC: yes
	Age: 46.0 (range = 21 to 86)
	Gender (per cent men): 67.8%
	Severity: overall score not reported
	INCLUSION CRITERIA
	• Plaques psoriasis in remission after 3 to 4 weeks treatment with Betamethasone
	dipropionate (erythema score = 1; induration = 0.5; scaling = 0)
	Note: 94/123 (76%) achieved remission during acute phase
	EXCLUSION CRITERIA
	• Recent topical or systemic treatment
	• Pregnant
	Nursing
	Intent to conceive
	• Not achieving remission during acute phase treatment

Katz 1991a (Continued)

Interventions	 Betamethasone dipropionate, intermittent maintenance (3 doses at 12-hour intervals once a week) (B) Placebo (vehicle) (P)
Outcomes	 Signs (erythema; induration; scaling) TSS (0 to 9) Area adjusted clinical score Treatment failure (adjusted clinical score ≥ 2.5, or overall disease status moderate or severe) Overall disease status Patient evaluation of effectiveness. Time to relapse
Notes	The trial did not report sponsorship, but the Schering Corporation employed the cor- responding author

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	4.3%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Katz 1991b

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: identical tubes Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	labelled by computer-generated code
Participants	N: 110 Treatment duration: 2 wks LF: 2 (1.8%) BC: inadequately reported Age: 51.7 (range = 19 to 84) Gender (per cent men): 80.6% Severity: Total Severity Score (0 to 12) (median) = 8 Duration of disease (years): 21 (range 1 to 57) Duration of exacerbation (years): 15.4 (range < 1 to 57) INCLUSION CRITERIA • Comparable bilateral lesions of moderate or greater severity of plaque psoriasis • Adult • At least 2 signs or symptoms of at least moderate severity • Lesions ≥ 10 cm ² EXCLUSION CRITERIA • Pustular or erythodermic psoriasis • Recent topical or systemic medication • Women at risk of pregnancy	
Interventions	 Halobetasol propionate cream 0.05% BD (H) Placebo (vehicle) (P) 	
Outcomes	 Severity (0 to 3) (erythema; plaque elevation; scaling pruritis) Total Severity Score (0 to 12) Patient Global Assessments of effectiveness and overall rating (5-pt: poor to excellent) 	
Notes	The trial was supported by an educational grant from Westwood-Squibb Pharmaceuti- cals, a Bristol-Myers Squibb company	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Katz 1991b (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	1.8%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Kaufmann 2002 (H)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation schedule Concealment: unclear BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described
Participants	 N: 1603 Treatment duration: 4 wks; FU: 4 wks LF: 0 (0%) BC: yes Age: 48.4 (range = 17 to 90) Gender (per cent men): 60.5% Severity: PASI mean = 10.0 (range = 1.2 to 49.5) Duration: 19.2 (range = 0 to 75) INCLUSION CRITERIA People aged 18 and over with chronic plaque psoriasis BSA at least 10% EXCLUSION CRITERIA Unstable psoriasis in treatment areas Other skin diseases that could confound treatment assessments Concomitant antipsoriatic therapy Hypercalcaemia Application of study corticosteroid to untargeted lesion Pregnancy Lactation
Interventions	• Calcipotriol 50 mcg/g + betamethasone dipropionate 0.5 mg/g combination ointment OD (D)

Kaufmann 2002 (H) (Continued)

	 Calcipotriol 50 mcg/g, in combination vehicle ointment OD (C) Betamethasone dipropionate 0.5 mg/g, in combination vehicle ointment OD (B) Placebo (combination vehicle) ointment OD (P)
Outcomes	 PASI, modified (change score) Investigator's Global Assessment of Disease Severity (6-pt: disease absent to very severe) Patient's global assessment of disease severity (6-pt: worse to cleared)
Notes	Leo Pharmaceuticals sponsored the trial. Compliance rates were reported for each regimen.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Kaufmann 2002 (P)

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated randomisation schedule
	Concealment: unclear
	BLINDING
	Double-blind (participant/assessor)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 1603
1 articipanto	Treatment duration: 4 wks; FU: 4 wks
	LF: 0 (0%)
	BC: yes

Kaufmann 2002 (P) (Continued)

	Age: 48.4 (range = 17 to 90)
	Gender (per cent men): 60.5%
	Severity: PASI mean = 10.0 (range = 1.2 to 49.5)
	Duration: 19.2 (range = 0 to 75)
	INCLUSION CRITERIA
	• People aged 18 and over with chronic plaque psoriasis
	• BSA at least 10%
	EXCLUSION CRITERIA
	• Unstable psoriasis in treatment areas
	• Other skin diseases that could confound treatment assessments
	• Concomitant antipsoriatic therapy
	• Hypercalcaemia
	• Application of study corticosteroid to untargeted lesion
	• Pregnancy
	• Lactation
Interventions	 Calcipotriol 50 mcg/g + betamethasone dipropionate 0.5 mg/g combination ointment, OD (D) Calcipotriol 50 mcg/g, in combination vehicle ointment OD (C) Betamethasone dipropionate 0.5 mg/g, in combination vehicle ointment OD (B) Placebo (combination vehicle) ointment OD (P)
Outcomes	 PASI, modified (change score) Investigator's Global Assessment of Disease Severity (6-pt: disease absent to very severe) Patient's global assessment of disease severity (6-pt: worse to cleared)
Notes	Leo Pharmaceuticals sponsored the trial. Compliance were rates reported for each regimen.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Kim 1994

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (unclear who was blinded) WITHDRAWAL/DROPOUT Not described	
Participants	N: 10 Treatment duration: 8 wks; FU: 8 wks LF: 0 (0%) BC: yes Age: 32.1 (range 20 to 52) Gender (per cent men): 60% Severity: PASI = 10.49 (1.60SD) Duration: 6.7 years (range 0.2 to 15) INCLUSION CRITERIA • Psoriasis EXCLUSION CRITERIA • Not identifiable	
Interventions	 Calcipotriol ointment 50 mcg/g BD (Desoxymethasone ointment 2.5 mg/g 	
Outcomes	 PASI Erythema, infiltration, desquamation 	(0 to 9)
Notes	The trial did not report sponsorship.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (unclear who was blinded).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were reported.

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Kim 1994 (Continued)

Baseline comparability demonstrated	Low risk	-
iss 1996		
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not r Concealment: unclear BLINDING Double-blind (participant/inves WITHDRAWAL/DROPOUT Described	
Participants	N: 239 Treatment duration: 8 wks LF: 29 (12.1%) BC: not reported Age: not reported Gender (per cent men): not reported INCLUSION CRITERIA • Moderate scalp psoriasis • Adult • Overall disease severity ≥ 4 EXCLUSION CRITERIA • None reported	
Interventions	 Calcipotriene solution 0.0025% and 0.005% BD (C) Placebo (vehicle) (P) 	
Outcomes	 Severity (scaling; erythema; plaque elevation; pruritis) Overall severity (9-pt: none to very severe) Investigator Global Assessment (4-pt: worsened to cleared) 	
Notes	Bristol Myers Squibb Pharmaceuticals sponsored the trial. Carder 1996 reports finding for subgroup (N = 29). This was a scalp trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Kiss 1996 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	12.1%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Klaber 1994

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: Concealment: unclear BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described
Participants	N: 474 Treatment duration: 4 wks LF: assessment = 6 (1.3%) TSS: 29 (6.1%) BC: yes Age: 44.1 (range = 18 to 90) Gender (per cent men): 51.5% Severity: TSS (0 to 12) = 6.5 (range = 2 to 12) Duration of scalp psoriasis (yrs): 13.1 (range = 0.1 to 67.0) INCLUSION CRITERIA • Adults • Adults • Stable, mild-to-moderate scalp psoriasis • History of psoriasis on body EXCLUSION CRITERIA • More extensive, severe, or infected psoriasis • Recent systemic antipsoriatic treatment or UV • Concurrent vitamin D, calcium, or other relevant medication • Significant hepatic or renal disease • Hypercalcaemia • Risk of pregnancy • Pregnancy • Lactation

Klaber 1994 (Continued)

Interventions	 Calcipotriol solution 50 mcg/ml BD (C) Betamethasone 17-valerate solution 1 mg/ml BD (B)
Outcomes	 Investigator Global Assessment (5-pt: worse to cleared) Patient Global Assessment (5-pt: worse to cleared) Total Sign Score (erythema, thickness, scaliness) (0 to 12) Assessment of extent of scalp psoriasis Assessment of acceptability
Notes	Leo Pharmaceutical Products sponsored the trial. This was a scalp trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	1.3%
Baseline assessments	Low risk	reported
Baseline comparability demonstrated	Low risk	-

Klaber 2000b

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Open
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 475 Treatment duration: 8 wks; FU: 24 wks (N = 166) LF: 52 (10.9%) BC: yes

	Age: 45.3
	Gender (per cent men): 52.0%
	Severity: Total Severity Score (0 to 12) = 5.1
	INCLUSION CRITERIA
	Mild or moderate scalp psoriasis
	EXCLUSION CRITERIA
	• Other forms of psoriasis
	• Topical antipsoriatic treatment within previous 2 wks
	• Systemic antipsoriatic treatment or UV therapy within previous 4 wks
	• Concomitant vitamins, calcium, or other medications that could affect the course
	of psoriasis
	Known hypersensitivity to study medications
	• Pregnancy
	Inadequate contraception
	Lactation
	• Hypercalaemia
	Significant renal or hepatic disease
T	
Interventions	• Calcipotriol solution 50 mcg/g (Dovonex®) BD (C)
	• Coal tar 1%, coconut oil 1%, salicylic acid 0.5%, shampoo (Capasal ®) OD (T)
Outcomes	1. IAGI (6-pt: worse to cleared)
Outcomes	2. TSS (0 to 12)
	3. Patients global assessment of disease severity (VAS)
Notes	Leo Pharmaceuticals sponsored the trial.
	All participants who achieved at least a slight improvement in scalp psoriasis then received
	16 weeks of treatment with calcipotriol BD.
	This was a scalp trial.
	I

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	10.9%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Koo 2006

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated (1:1:2) Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 86 Treatment duration: 2 wks; FU: 2 wks LF: 0 (0%) BC: not demonstrated Age: NR Gender (per cent men): NR Severity: NR INCLUSION CRITERIA People aged > = 18 with plaque psoriasis Good general health Target lesions on trunk and extremities with severity score > = 2 for each sign (erythema, induration, scaling; 0 to 4) EXCLUSION CRITERIA Pregnancy or risk thereof Lactation Use of investigational drug within previous 30 dys Systemic or phototherapy within previous 30 dys Use of topical or intralesional therapies (other than emollients) within previous 2 wks Guttate, erythrodermic, pustular psoriasis BSA > 20% (excluding face/scalp) Known hypersensitivity to study medications Concomitant topical/systemic medication that might affect target lesions Concomitant phototherapy
Interventions	 Part 1 Calcipotriene 0.005% ointment BD (C) Clobetasol propionate foam 0.05% BD (CP) Calcipotriene 0.005% ointment plus clobetasol propionate foam 0.05% BD (CC) Part 2 Follow-up maintenance study based on responders from combination group (CC)
Outcomes	 Total Severity Score, based on Psoriasis Grading Scale (erythema, induration, scaling, each 0 to 4) (TSS; 0 to 12). Scores adjusted for baseline differences. TSS reported separately for trunk and extremity lesions Remission: per cent reduction in TSS > = 50%, IGA reduction of 3 points, or PGA reduction of 3 points Investigator Global Severity Assessment (IGA); 7-pt: 0 (clear) to 6 (severe)

Koo 2006 (Continued)

	 4. Subject Global Severity Assessment (PGA); 7-pt: 0 (clear) to 6 (severe) 5. Adverse events 6. Skin atrophy 7. Safety
Notes	Connetics Corporation, Palo Alto, California, US, sponsored the trial We sought data, but we received no response. There was SD imputation (IGA/PGA).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	This was not demonstrated.

Kragballe 1988b

Methods	DESIGN Within-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Not described
Participants	N: 30
	Treatment duration: 6 wks
	LF: 3 (10%)
	BC: yes
	Age: $39 (range = 18 \text{ to } 65)$
	Gender (per cent men): 30.0%
	Severity: TSS (0 to 9) = 6.9; per cent BSA = 18.7% (range = 12% to 50%)

Duration (yrs): 15.3 (range = 1 to 35)
INCLUSION CRITERIA
Stable symmetrically distributed moderate
Chronic plaque-type psoriasis
• Outpatients
• Adult
Women above child bearing age or using adequate contraception
EXCLUSION CRITERIA
• Recent topical, systemic, intralesional, or UV radiation therapy (excluding bland
emollients)
 Non-normal serum levels of calcium and creatinine
Taking calcium tablets
• Calcipotriol cream 10 mcg/g, 33 mcg/g, or 100 mcg/g BD C (10); C (33); C (100)
• Placebo (vehicle) (P)
1. Severity (erythema; thickness; scaling)
2. TSS (0 to 9)
3. Investigator Global Assessment (5-pt: worse to clear)
4. Patient Global Assessment (5-pt: worse to clear)
The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	10.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Kragballe 1991a

Methods	DESIGN
	Within-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not stated
	Concealment: unclear
	BLINDING
	Double-blind (participant/assessor)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 345
-	Treatment duration: 6 wks
	LF: 3 (0.9%)
	BC: yes
	Age: 45.2 (range = 18 to 90)
	Gender (per cent men): 58.8%
	Severity: PASI = 8.35 (range = 0.60 to 48.5)
	Duration (yrs): 19.5 (range = 0.5 to 76)
	INCLUSION CRITERIA
	• Adult
	Symmetrical chronic plaque psoriasis
	Inpatients and outpatients
	EXCLUSION CRITERIA
	• Unstable psoriasis
	Recent systemic or UV therapy
	• Hypercalcaemia
	Impaired renal/hepatic function High dags solaium (Vitamin D inteles
	High dose calcium/Vitamin D intakeUnresponsive to corticosteroids
	Concomitant medication
Interventions	• Calcipotriol ointment 50 mcg/g BD (C)
	• Betamethasone valerate ointment 0.1% BD (B)
Outcomes	1. PASI
Cuttonits	 Total Sign Score (erythema, thickness, scaliness) (0 to 12)
	3. Patient assessment of response
Notes	Leo Pharmaceuticals sponsored the trial
	Trial participants were inpatients and outpatients.
	There was SD imputation (TSS).

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Kragballe 1991a (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.9%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Kragballe 1998b

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: not stated BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described
Participants	 N: 699 Treatment duration: 8 wks; FU: 8 wks LF: 8 (1.1%) BC: psoriasis comparable, demographics unclear Age: not stated Gender (per cent men): not stated Severity: not stated INCLUSION CRITERIA Adult Stable chronic plaque psoriasis on trunk and limbs EXCLUSION CRITERIA Pregnancy Risk of pregnancy Lactation Recent systemic or UV therapy Concomitant medication Hypercalcaemia or renal disease Planned exposure to sun
Interventions	 Calcipotriol cream 50 mcg/g BD (C2) Calcipotriol cream 50 mcg/g OM, plus clobetasone17-butyrate cream,0.5 mg/g ON (CL)

Kragballe 1998b (Continued)

	 Calcipotriol cream 50 mcg/g OM, plus betamethasone 17-valerate cream 1 mg/g ON (CB) Calcipotriol cream 50 mcg/g OM, plus vehicle ON (C1)
Outcomes	 PASI Investigator overall assessment of response (6-pt: worse to clearance) Patient overall assessment of response (6-pt: worse to clearance)
Notes	Leo Pharmaceuticals sponsored the trial

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The study was double-blind (participant/ assessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	1.1%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Kragballe 2004

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated randomisation schedule, using cen-
	tralised telephone voice response system
	Concealment: adequate
	BLINDING
	Double-blind (participant/investigator) (Groups A and B)
	Single-blind (investigator) (Group C)
	WITHDRAWAL/DROPOUT
	Described
D	NL 072
Participants	N: 972
	Treatment duration: 8 wk; FU: 12 wks
	LF: 99 (10.2%)
	BC: yes

Kragballe 2004 (Continued)

	Age: 47.7 (range = 18 to 97)	
	Gender (per cent men): 63.8%	
	Severity: PASI = 10.5 (range = 2 to 49); per cent with moderate disease = 64.3%	
	Duration (yrs): 18.5 (range = 0 to 70)	
	INCLUSION CRITERIA	
	• Aged 18 and over	
	Chronic plaque psoriasis amenable to topical treatment	
	 BSA ≥ 10% of at least 1 body region (arms, trunk, legs) 	
	EXCLUSION CRITERIA	
	• Pregnancy or risk thereof	
	Lactation	
	• Unstable psoriasis or other inflammatory skin disease	
	• Concurrent systemic or UV therapy	
	• Concurrent topical therapy for trunk or limbs	
	Abnormal calcium homeostasis	
Interventions	 TCP OD for 8 wks then calcipotriol ointment 50 mcg/g OD for 4 wks (A) TCP OD for 4 wks then calcipotriol ointment 50 mcg/g OD (weekdays) and TCP OD (weekends) for 8 wks (B) Calcipotriol ointment 50 mcg/g BD for 12 wks (C) TCP: calcipotriol ointment 50 mcg/g, plus betamethasone dipropionate 0.5 mg/g ointment 	
Outcomes	 PASI Investigator's global assessment of severity (PGA) (6-pt: absence of disease to very severe disease) Self-reported compliance with trial medication 	
Notes	Leo Pharmaceuticals sponsored the trial. Request data: none supplied Reversible skin atrophy: A: 1/322; B: 0/322; C: 0/327	

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	A centralised telephone voice response sys- tem was used for the participant assign- ment, and this removed the opportunity for investigator bias during randomisation
Blinding (performance bias and detection bias) All outcomes	Low risk	Groups A and B were double-blind (par- ticipant/investigator); group C was single- blind (investigator)
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	10.2%

Kragballe 2004 (Continued)

Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Kragballe 2006

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation schedule (1:1:1) Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 634 Treatment duration: 52 wks; FU: 52 wks LF: 8 (1.3%) BC: yes Age: 48.7 (14.2SD) Gender (per cent men): 61.0% Ethnicity (per cent white): 97.3% Severity: per cent IGA moderate = 69.1%; per cent IGA severe = 27.9%; per cent IGA very severe = 3.0% Duration (yrs): mean = 19.7; median = 17.0; range = 1 to 65 Use of topical corticosteroids during last decade (mths): 33.8 INCLUSION CRITERIA • Outpatients aged > = 18 with plaque psoriasis of trunk and limbs • IGA at least moderate. EXCLUSION CRITERIA • Erythrodermic, exfoliative, and pustular psoriasis • Skin infection, use of systemic or topical antipsoriatic therapy • Use of PUVA or UVB • BSA > = 30% (and requiring treatment) • Calcium metabolic disorder • Pregnancy • Lactation
Interventions	 Calcipotriol ointment 50 mcg/g plus betamethasone dipropionate ointment 0.5 mg/g OD (C-B) Alternating every 4 weeks: calcipotriol ointment 50 mcg/g plus betamethasone dipropionate ointment 0.5 mg/g OD (4 wks) then calcipotriol 50 mcg/g (4 wks) (A) Calcipotriol ointment 50 mcg/g plus betamethasone dipropionate ointment 0.5 mg/g, OD (4 wks) then calcipotriol ointment 50 mcg/g (48 wks) (C) < = 100 g/wk

Kragballe 2006 (Continued)

Outcomes	 Investigator's Global Assessment (IGA): 6-pt (absent, very mild, mild, moderate, severe, very severe) Treatment success: IGA absent, very mild, or mild Independent assessment of adverse events by non-investigator clinicians: adjudicated corticosteroid reactions Patient Global Assessment (PGA): 3-pt (satisfactory, not satisfactory, not applicable (not used)) Compliance (medication usage) HPA assessed in subgroup of 19 participants
Notes	Leo Pharma A/S sponsored the trial. The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	A computer-generated randomisation schedule was used (1:1:1)
Loss to follow up	Low risk	1.3%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Unclear risk	The trial only reported mean values (no measure of spread). They described groups as "well balanced"

Kragballe 2009

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated schedule (1:2) Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 312 Treatment duration: 8 wks; FU: 16 wks LF: 2 (0.6%) BC: yes Age: 51.0 (15.4SD); range = 18 to 91 Gender (per cent men): 43.0% Ethnicity (per cent white): 99.0% Severity: IGA moderate = 56.7%; IGA severe = 35.9%; IGA very severe = 7.4%; TSS (0 to 12) = 7.3 (1.7SD) Duration (yrs): 18.7 (14.6SD); range = 0 to 70 INCLUSION CRITERIA • People with moderately severe scalp psoriasis amenable to treatment with < = 100 g medication/wk (combined product) or 60 mL solution/wk (calcipotriol) • Psoriasis of trunk/limbs • Scalp involvement > = 10% • Clinical signs > = moderate for > = 1 sign and > = slight for remaining signs • IGA > = moderate EXCLUSION CRITERIA • PUVA or grenz ray therapy within previous 4 wks • Topical scalp treatments or UVB therapy or very potent (WHO group IV) corticosteroids on body within previous 2 wks • Biologicals within previous 6 mths • Systemic therapies within previous 4 wks • Current diagnosis of unstable forms of psoriasis or other skin diseases confounding psoriasis assessment • Skin infections • Atrophy of the scalp • Calcium homeostasis abnormality • Severe renal or hepatic disorder • Concomitant use of medications that could affect scalp psoriasis • Pregnancy • Lactation
Interventions	 Calcipotriol solution 50 mcg/g BD (C) Calcipotriol gel 50 mcg/g plus betamethasone dipropionate "scalp formulation" (gel) 0.5 mg/g OD (C-B) There was no concomitant topical treatment, emollient, or medicated shampoo or conditioner

Kragballe 2009 (Continued)

Outcomes	 Investigator's Global Assessment (IGA): 6-pt (absent (0), very mild, mild, moderate, severe, very severe (5)) Treatment success: IGA clear or minimal Total Sign Score (TSS): 0 to 12 Patient Global Assessment of Disease Severity (PGA). 5-pt scale (clear, very mild, mild, moderate, severe) Detions assessment of inshing (4 ptr page mild moderate severe)
	 5. Patient assessment of itching (4-pt: none, mild, moderate, severe) 6. Compliance: self-report; use of study medication
Notes	Leo Pharma A/S, Ballerup, Denmark, sponsored the trial. The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	A computer-generated randomisation schedule was used (1:2).
Loss to follow up	Low risk	0.6%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Kreuter 2006 (H)

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated randomisation lists
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 80 Treatment duration: 4 wks; FU: 10 wks LF: 2 (2.5%)

Kreuter 2006 (H) (Continued)

Age: 52.4 (14.3SD); range = 28 to 87 Gender (per cent men): 61.3% Ethnicity: NR Severity: mPASI = 21.2 (13.4SD); range = 3 to 54; VAS (0 to 10) = 4.8 (2.7SD); ra = 1 to 10 INCLUSION CRITERIA • People aged > = 18 with inverse psoriasis that had been present for at least 6 m
Gender (per cent men): 61.3% Ethnicity: NR Severity: mPASI = 21.2 (13.4SD); range = 3 to 54; VAS (0 to 10) = 4.8 (2.7SD); ra = 1 to 10 INCLUSION CRITERIA • People aged > = 18 with inverse psoriasis that had been present for at least 6 m
Ethnicity: NR Severity: mPASI = 21.2 (13.4SD); range = 3 to 54; VAS (0 to 10) = 4.8 (2.7SD); ra = 1 to 10 INCLUSION CRITERIA • People aged > = 18 with inverse psoriasis that had been present for at least 6 m
Severity: mPASI = 21.2 (13.4SD); range = 3 to 54; VAS (0 to 10) = 4.8 (2.7SD); ra = 1 to 10 INCLUSION CRITERIA • People aged > = 18 with inverse psoriasis that had been present for at least 6 m
 = 1 to 10 INCLUSION CRITERIA People aged > = 18 with inverse psoriasis that had been present for at least 6 m
• People aged > = 18 with inverse psoriasis that had been present for at least 6 m
Good general health
EXCLUSION CRITERIA
• Systemic or phototherapies within previous 4 wks
 Topical therapies within previous 2 wks
Acute guttate or pustular psoriasis
• Pregnancy
• Lactation
 Severe concurrent infectious diseases
• Diseases associated with immunesuppression or malignancy
• Drug dependency
Mental dysfunction or other factors limiting compliance
Interventions • Calcipotriol 0.005% ointment OD (C)
• Betamethasone valerate 0.1% OD (B)
• Pimecrolimus 1% cream OD (PM)
• Placebo (vehicle for pimecrolimus) OD (P)
No concomitant therapies were permitted (including emollients)
Outcomes1. mPASI (0 to 72): assessment confined to body area affected. Each sign scored (4 and then multiplied by area affects (0 (0%) to 6 (90% to 100%))
 VAS (itching): 0 = absence to 10 = maximum Adverse events
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3. Adverse events
3. Adverse events Notes Novartis Pharma GmbH sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Computer-generated randomisation lists were used.

Kreuter 2006 (H) (Continued)

Loss to follow up	Low risk	2.5%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Kreuter 2006 (P)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation lists Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 80 Treatment duration: 4 wks; FU: 10 wks LF: 2 (2.5%) BC: yes Age: 52.4 (14.3SD); range = 28 to 87 Gender (per cent men): 61.3% Ethnicity: NR Severity: mPASI = 21.2 (13.4SD), range = 3 to 54; VAS (0 to 10) = 4.8 (2.7SD), range = 1 to 10 INCLUSION CRITERIA • People aged > = 18 with inverse psoriasis that had been present for at least 6 mths • Good general health EXCLUSION CRITERIA • Systemic or phototherapies within previous 4 wks • Topical therapies within previous 2 wks • Acute guttate or pustular psoriasis • Pregnancy • Lactation • Severe concurrent infectious diseases • Diseases associated with immunesuppression or malignancy • Drug dependency • Mental dysfunction or other factors limiting compliance
Interventions	 Calcipotriol 0.005% ointment OD (C) Betamethasone valerate 0.1% OD (B) Pimecrolimus 1% cream OD (PM) Placebo (vehicle for pimecrolimus) OD (P) No concomitant therapies were permitted (including emollients)

Kreuter 2006 (P) (Continued)

Outcomes	 mPASI (0 to 72): assessment confined to body area affected. Each sign scored 0 to 4 and then multiplied by area affects (0 (0%) to 6 (90% to 100%)) VAS (itching): 0 = absence to 10 = maximum Adverse events
Notes	Novartis Pharma GmbH sponsored the trial. Atrophy was not reported. We sought unpublished data, but it was reported to be unavailable There was SD imputation (PASI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Computer-generated randomisation lists were used.
Loss to follow up	Low risk	2.5%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Krueger 1998

Methods	DESIGN Within-patient
	Participant delivery ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 45
1	Treatment duration: 6 wks
	LF: 0 (0%)
	BC: reported to be similar
	Age: $50 (range = 23 \text{ to } 83)$
	Age: $50 (tange = 25 to 65)$

Krueger 1998 (Continued)

	 Gender (per cent men): 84% Severity: not reported INCLUSION CRITERIA Mild to moderate bilateral psoriatic plaques Adult Total Severity Score less than or equal to 6. EXCLUSION CRITERIA Pregnant Nursing or of likely to conceive Recent use of certain topical agents Recent systemic retinoids, UV phototherapy, or systemic antipsoriasis drugs
Interventions	 Tazarotene gel 0.01% or 0.05% BD (T) Placebo (vehicle) (P)
Outcomes	 Severity (erythema; plaque elevation; scaling) TSS (0 to 12) Investigator Global Assessment (6-pt: no change/worse to completely clear)
Notes	Allergan, Inc, California, sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Unclear risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	These were stated to be similar; compara- bility was not demonstrated

Köse 1997

Methods	DESIGN Between-patient Delivery unclear ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Unclear WITHDRAWAL/DROPOUT Described
Participants	N: 43 Treatment duration: 10 days FU: 20 days LF: 0 (0%) BC: yes Age: not stated Gender (per cent men): not stated Severity: not stated INCLUSION CRITERIA • Psoriasis of the scalp EXCLUSION CRITERIA • Not reported
Interventions	 Calcipotriol ointment 50 mcg/g occluded ON (CO) Clobetasol 17-propionate solution BD (CP)
Outcomes	1. TSS (0 to 9)
Notes	The trial did not report sponsorship. This was a scalp trial.
Risk of hias	

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Unclear risk	The trial did not report these.

Köse 1997 (Continued)

Baseline comparability demonstrated	Unclear risk	The trial reported baseline comparabil- ity (demographic/clinical), but it was not demonstrated
Lahfa 2003		
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described	
Participants		
Interventions	 Calcipotriol 50 mcg/g ointment ON, plus clobetasol propionate 0.05% cream OM (2 to 4 wks), then calcipotriol ointment BD (to wk 12) (C1) Calcitriol 3 mcg/g ointment ON, plus clobetasol propionate 0.05% cream OM (2 to 4 wks), then calcitriol ointment BD (to wk 12) (C2) In the initial phase, the participant applied dual therapy until achieving clearance or marked improvement or until 4 wks elapsed, then switched to monotherapy maintenance phase 	

Lahfa 2003 (Continued)

Outcomes	 PASI, based on scores from 3 lesions, 1 each on trunk, and upper and lower limbs Investigator's Assessment of Global Improvement (IAGI): 7-pt (worse to clear, rescaled 0 to 6) Clinical success: IAGI > = marked improved Relapse (maintenance phase): IAGI < = moderate improvement Patient Assessment of Global Improvement: 7-pt (worse to clear) Adverse events: cutaneous safety including signs and soreness (all scored 0 to 3); investigator assessment of global safety (poor, good, excellent); patient self report
Notes	Galderma Laboratories sponsored the trial. We sought unpublished data, but it was reported to be unavailable There was SD imputation (PASI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	4.0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Landi 1993

DESIGN
Between-patient
Delivery unclear
ALLOCATION
Random
Method of randomisation: not stated
Concealment: unclear
BLINDING
Unclear
WITHDRAWAL/DROPOUT
Described
N: 40 Treatment duration: 6 wks; FU: 10 wks LF: 0 (0%)

Landi 1993 (Continued)

	 BC: psoriasis comparable, demographics unclear Age: range = 17 to 84 Gender (per cent men): not stated Severity: PASI (mean) = 11.6 (range = 3.0 to 35.1) INCLUSION CRITERIA Adult Mild and moderate psoriasis EXCLUSION CRITERIA Not reported
Interventions	 Calcipotriol ointment 50 mcg/g BD (C) Clobetasol propionate 0.05% ointment BD (CP)
Outcomes	1. PASI
Notes	Leo Pharmaceuticals sponsored the trial. Landi 1993 reported the findings of a single centre, 1 of 3 centres reported in Landi 1993 (N = 120).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Unclear risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Lane 1983

Risk of bias	There was SD imputation (TSS).		
Outcomes		 Severity (scaling; erythema; pruritis; thickening; crusting; overall condition) Total Severity Score (0 to 20) 	
Interventions		 Betamethasone dipropionate ointment 0.05% OD (B) Diflorasone diacetate ointment 0.05% OD (D) Placebo (vehicle) (P) 	
Participants	LF: 18 (11.5%) BC: yes Age: 39.6 Gender (per cent men): 52.5% Severity: TSS (0 to 20) = 10.6 INCLUSION CRITERIA • History and physical finding erythema, epidermal thickening, • All ages > 1 year • Stable disease. EXCLUSION CRITERIA • Recent topical or systemic co • Oral antihistamine	 N: 157 Treatment duration: 3 wks; FU: 3 wks LF: 18 (11.5%) BC: yes Age: 39.6 Gender (per cent men): 52.5% Severity: TSS (0 to 20) = 10.6 INCLUSION CRITERIA History and physical finding compatible with psoriasis including scaling erythema, epidermal thickening, crusting, or both All ages > 1 year Stable disease. EXCLUSION CRITERIA Recent topical or systemic corticosteroid treatment Oral antihistamine Antipruritic therapy, UV, or X-ray therapy or any medication affecting the study 	
Methods	Between-patient Participant delivery ALLOCATION Random Method of randomisation: not re Concealment: unclear BLINDING	Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT	

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Lane 1983 (Continued)

Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	11.5%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Langley 2011 (H)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated (2:2:1) Concealment: adequate: treatments dispensed by non-study staff BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 458 Treatment duration: 8 wks; FU: 16 wks LF: 0 (0%) BC: yes Age: 51.6 (14SD) Gender (per cent men): 62.2% Ethnicity (per cent white): 93.9% Severity: mPASI = 9.39, range = 2.4 to 59.4; IGA moderate = 68.3%; IGA severe = 29. 5%; IGA very severe = 2.2%; per cent BSA = 9.3 (8.2SD) Duration (yrs): 19.8 (13.3SD) INCLUSION CRITERIA Participants with moderately severe plaque psoriasis involving trunk, limbs, or both BSA > = 10% on arms, trunk, limbs, or a mixture of the aforementioned Amendable to maximum weekly dose of 100 g of gel or 70 g of ointment EXCLUSION CRITERIA Systemic biological therapy within previous 3 mths Other systemic treatment within previous 4 wks Concurrent oral vitamin D > 500 IU/day UVA or grenz ray therapy within previous 4 wks UVB within previous 2 wks Pregnancy Lactation

Langley 2011 (H) (Continued)

Interventions	 Tacalcitol ointment 4 mcg/g OD (T) Calcipotriol gel 50 mcg/g plus betamethasone dipropionate gel 0.5 mg/g OD (C-B) Placebo (gel vehicle) OD (P) Participants achieving clear IGA stopped treatment and restarted if psoriasis reappeared There was follow up of responders only (IGA clear or almost clear)
Outcomes	 Investigator's Global Assessment of Disease Severity: 6-pt (clear to very severe) Treatment responder (controlled disease): IGA clear or almost clear at wk 8 mPASI (0 to 64.8) PASI 50, PASI 75 Patient Global Assessment (PGA) of disease severity: 5-pt (clear to severe) Adverse events Relapse (treatment responders only): > = 50% reduction in PASI improvement achieved (wk0 to wk8) Time to relapse Rebound (treatment responders only): > 125% reduction in PASI at wk 0 Compliance (wk 2 to wk 6): self-report and medication usage Adverse events
Notes	Leo Pharma A/S, Ballerup, Denmark, sponsored the trial. Participants could not be completely blinded to treatment allocation because of differ- ences in formulation and packaging (bottles for gel vs. tubes for ointments) Atrophy was not assessed. The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	A designated third person performed allo- cation of the investigational products at all 18 centres
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator); it was not possible to blind participants to treatments because of differences in vehicle formulations
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Langley 2011 (P)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated (2:2:1) Concealment: adequate: treatments dispensed by non-study staff. BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 458 Treatment duration: 8 wks; FU: 16 wks LF: 0 (0%) BC: yes Age: 51.6 (14SD) Gender (per cent men): 62.2% Ethnicity (per cent white): 93.9% Severity: mPASI = 9.39, range = 2.4 to 59.4; IGA moderate = 68.3%; IGA severe = 29. 5%; IGA very severe = 2.2%; per cent BSA = 9.3 (8.2SD) Duration (yrs): 19.8 (13.3SD) INCLUSION CRITERIA • Participants with moderately severe plaque psoriasis involving trunk, limbs, or both • BSA > = 10% on arms, trunk, limbs, or a mixture of the aforementioned • Amendable to maximum weekly dose of 100 g of gel or 70 g of ointment EXCLUSION CRITERIA • Systemic biological therapy within previous 3 mths • Other systemic treatment within previous 4 wks • Concurrent oral vitamin D > 500 IU/day • UVA or grenz ray therapy within previous 4 wks • UVB within previous 2 wks • Pregnancy • Lactation
Interventions	 Tacalcitol ointment 4 mcg/g OD (T) Calcipotriol gel 50 mcg/g plus betamethasone dipropionate gel 0.5 mg/g OD (C-B) Placebo (gel vehicle) OD (P) Participants achieving clear IGA stopped treatment and restarted if psoriasis reappeared There was follow up of responders only (IGA clear or almost clear)
Outcomes	 Investigator's Global Assessment of Disease Severity: 6-pt (clear to very severe) Treatment responder (controlled disease): IGA clear or almost clear at wk 8 mPASI (0 to 64.8) PASI 50, PASI 75 Patient Global Assessment (PGA) of disease severity: 5-pt (clear to severe) Adverse events Relapse (treatment responders only): > = 50% reduction in PASI improvement

Langley 2011 (P) (Continued)

	 achieved (wk0 to wk8) 8. Time to relapse 9. Rebound (treatment responders only): > 125% reduction in PASI at wk 0 10. Compliance (wk 2 to wk 6): self-report and medication usage 11. Adverse events
Notes	Leo Pharma A/S, Ballerup, Denmark, sponsored the trial. Participants could not be completely blinded to treatment allocation because of differ- ences in formulation and packaging (bottles for gel vs. tubes for ointments) Atrophy was not assessed. The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	A designated third person performed allo- cation of the investigational products at all 18 centres
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator); it was not possible to blind participants to treatments because of differences in vehicle formulations
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Langner 1992

Methods

DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described

Langner 1992 (Continued)

Participants	N: 29 Treatment duration: 6 wks; FU: 6 wks LF: 0 (0%) BC: yes Age: mean: 40.5 (range = 16-77) Gender (per cent men): 69.0% Severity: not reported INCLUSION CRITERIA • Severe chronic psoriasis • Symmetrical lesions • Adult • Outpatients EXCLUSION CRITERIA • Pregnancy or inadequate contraception
Interventions	 Calcitriol ointment 3 mcg/g BD (C) Placebo (vehicle) (P)
Outcomes	 Severity (erythema; pustules, desquamation, encrustation, vesiculation and pruritis) Investigator Global Assessment (6-pt: worse to clear)
Notes	The trial did not report sponsorship. All participants received 2 weeks' pre-treatment with vehicle BD

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Langner 1993

Methods	DESIGN
	Within-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: unclear
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 32
rarticipants	Treatment duration: 6 wks; FU: 6 wks
	LF: 2 (6.3%)
	BC: yes 42.4 (sense 16 to 77)
	Age: mean: 42.4 (range = 16 to 77)
	Gender (per cent men): 62.5%
	Severity: global severity score (0 to 4) = 3.5 INCLUSION CRITERIA
	Bilateral Summarial
	• Symmetrical
	Severe chronic plaque psoriasis
	Outpatients. EXCLUSION CRITERIA
	EXCLUSION CRITERIA
	Pregnancy or inadequate contraception
	• Use of calcium
	Vitamin D or analogues
	Calcium-containing antacids
	• Digitalis
	Thiazide diuretics or glucocorticosteroids
Interventions	• Calcitriol ointment 15 mcg/g BD (C)
	• Placebo (vehicle) (P)
2	
Outcomes	1. Severity (erythema; scaling; induration; pruritis)
	2. PASI
	3. Investigator Global Assessment (6-pt: worse to clear)
Notes	The trial did not report sponsorship.
	All participants received 2 weeks' pre-treatment with vehicle BD
Risk of bias	

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Langner 1993 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	6.3%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Langner 2001 (H)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 44 Treatment duration: 6 wks; FU: 6 wks (14 wks for responders) LF: 4 (9.1%) BC: not reported Age: not reported Gender (per cent men): 54.5% Severity: not reported INCLUSION CRITERIA • Adults with chronic plaque psoriasis • BSA = 20% EXCLUSION CRITERIA • Pregnancy • Inadequate contraception
Interventions	 Calcitriol ointment 3 mcg/g BD (C) Betamethasone valerate ointment 0.1% BD (B)
Outcomes	1. IAGI (5-pt: worse to clearance)
Notes	The trial did not report sponsorship. All participants received 2 weeks' pre-treatment with vehicle BD

Langner 2001 (H) (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	9.1%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Langner 2001 (P)

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: unclear
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Dentitierente	N: 44
Participants	
	Treatment duration: 6 wks; FU: 6 wks (14 wks for responders) LF: 4 (9.1%)
	BC: not reported
	Age: not reported
	Gender (per cent men): 54.5%
	Severity: not reported
	INCLUSION CRITERIA
	Adults with chronic plaque psoriasis
	• BSA = 20%
	EXCLUSION CRITERIA
	• Pregnancy
	Inadequate contraception

Langner 2001 (P) (Continued)

Interventions	 Calcitriol ointment 3 mcg/g BD (C) Placebo ointment BD (P)
Outcomes	1. IAGI (5-pt: worse to clearance)
Notes	The trial did not report sponsorship. All participants received 2 weeks' pre-treatment with vehicle BD

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	9.1%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Lassus 1991

Methods	DESIGN
Withous	
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: block randomisation
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 50
rarticipants	
	Treatment duration: 12 wks; FU: 12 wks
	LF: 8 (16%)
	BC: yes
	Age: 42.8 (range = 22 to 50; N = 42)
	Gender (per cent men): 45.2% (N = 42)
	Severity: TSS (0 to 12) = 7.72
	INCLUSION CRITERIA

Lassus 1991 (Continued)

	 Stable psoriasis of at least 1 years' duration Mild to moderate plaque psoriasis Nummular, discoid, or guttate psoriasis Stable aged 18 to 50 Localised lesions EXCLUSION CRITERIA Pregnancy Lactation Antipsoriatic therapy within previous 2 wks People declining to abstain from alcohol during treatment period
Interventions	 Oleum horwathiensis (Psoricur®) OD (O) Placebo OD (P)
Outcomes	 TSS (0 to 12) Severity (scaling, pruritis, erythema, induration) IAGI (5-pt: poor to healed)
Notes	The trial did not report sponsorship. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Block randomisation was used.
Loss to follow up	Low risk	16.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Lebwohl 2002

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported (but ratio 3:1 used)
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 81
i ii ticipiinto	Treatment duration: 2 wks; FU: 4 wks
	LF: 5 (6.2%)
	BC: yes
	Age: not reported
	Gender (per cent men): not reported
	Severity: pruritis $(0 \text{ to } 4) = 2.11$
	INCLUSION CRITERIA
	Mild to moderate plaque type psoriasis
	• Aged at least 18
	• TSS (0 to 12) ≥ 3
	• Target lesions in at least 1 of 5 anatomical regions
	• BSA $\leq 20\%$
	EXCLUSION CRITERIA
	 Investigational medication within previous 4 wks
	 Topical antipsoriatic treatment within previous 2 wks
	 Systemic antipsoriatic treatment within previous 4 wks
	 Concurrent UV treatment or sunbathing
	• Pregnancy
	• Lactation
	Inadequate contraception
	 Men wishing to father children during the study
	Concurrent drug or alcohol abuse
Interventions	• Clobetasol propionate foam 0.05% BD (maximum of 50 g/wk) (C)
Inter ventions	 Clobelasti propionale toani 0.05% BD (maximum of 50 g/wk) (C) Placebo foam BD (P)
	· · · · · · · · · · · · · · · · · · ·
Outcomes	1. IAGI (7-pt: worse to completely clear)
	2. PAGI (7-pt: worse to completely clear)
	3. Total Severity Score: erythema, scaling, thickness, pruritis (0 to 4)
	4. Adverse events
	5. Medicines consumption (compliance)
Notes	Connetics Corporation sponsored the trial.
INDICS	Only non-scalp sites were treated.

Lebwohl 2002 (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	6.2%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Lebwohl 2004

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 167
1 articipanto	Treatment duration: 8 wks; FU: 8 wks
	LF: 30 (18%)
	BC: yes
	Age: 48.0
	Gender (per cent men): 58.7%
	Severity: Static Severity Score (SSS) (0 to 6) (median) = 3 (range = 1.5 to 5.0); per cent
	with concurrent plaque-type lesions = 85%
	INCLUSION CRITERIA
	• Age limit unclear (stated as ≥ 2 and ≥ 16)
	 Age mine uneven (stated as <u>2</u> 2 and <u>2</u> 10) Chronic plaque psoriasis affecting intertriginous and facial skin
	• Disease stable or slowly worsening for ≥ 1 wk
	 Disease stable of slowly worsching for ≤ 1 wk Target lesion of moderate erythema and TSS (0 to 12) ≥ 4
	• Target residu of moderate crytitema and $135(0, 10, 12) \ge 4$ EXCLUSION CRITERIA
	Systemic therapy or phototherapy within previous 4 wks
	 Systemic therapy of photomerapy within previous 4 wks Topical therapy within previous 2 wks
	 Inhaled/intranasal corticosteroids within previous 2 wks
	 Other topical agents (excluding sunscreen) within previous 1 dy
	• Other topical agents (excluding subscieen) within previous 1 dy

Lebwohl 2004 (Continued)

	 Recently diagnosed (< 6 mths) or recent exacerbation of inverse psoriasis Uncontrolled chronic co-morbidity Pregnancy Lactation Previous use of tacrolimus ointment for facial or intertriginous psoriasis
Interventions	Tacrolimus ointment 0.1% BDPlacebo ointment BD
Outcomes	 Inverse psoriasis severity score (Static Severity Score) (SSS) (6-pt: clear to very severe) IAGI ('PGA') (7-pt: exacerbation to clear) Signs (0 to 3 each) (erythema, induration, desquamation; overall severity) Patient satisfaction (per cent agreeing with range of statements)
Notes	Fujisawa Healthcare Inc sponsored the trial. Loss to follow up was reported as 11/167. Non-study body sites received usual topical treatment. No adequate effectiveness data were reported or available from the sponsor. This was an inverse psoriasis trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	18.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Lebwohl 2007

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT	
Participants	Not described N: 418 Treatment duration: 8 wks; FU: 8 wks LF: 0 (0%) BC: NR separately (see notes) Age: NR Gender (per cent men): NR Severity: GSS (0 to 5) = 2.81 (0.40SD); DSS (0 to 12) (bony areas) = 6.4 (1.24SD); (non-bony areas) = 6.4 (1.33SD) INCLUSION CRITERIA • People aged > = 12 with mild to moderate plaque psoriasis • BSA <= 35% EXCLUSION CRITERIA • Not stated	
Interventions	 Calcitriol 3 mcg/g ointment BD (C) Placebo ointment BD (P) 	
Outcomes	 Global Severity Score (GSS) on a 6-pt scale (0 = clear, 1 = almost clear, 2 = mild, 3 moderate, 4 = severe, 5 = very severe) Dermatological sum score (DSS) (0 to 12): reported separately for bony and non- bony areas Investigator Assessment of Global Improvement (IAGI) on a 7-pt scale (clear to worse) Subject Global Assessment (PAGI), 7-pt (clear to worse) Routine clinical and safety laboratory parameters including calcium homeostasis (> 10% above upper limit of normal range) 	
Notes	Galderma R&D Inc, Princeton, New Jersey, sponsored the trial The main publication reports only pooled results and baseline statistics from 2 studies (Lebwohl 2007; Powers 2005). The sponsor supplied unpublished data.	
Risk of bias		
Bias	Authors' judgement	Support for judgement

Allocation concealment (selection bias)

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Unclear risk

The trial reported insufficient details.

Lebwohl 2007 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	This was not demonstrated.

Lee 2007

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 142 Treatment duration: 6 wks; FU: 14 wks LF: 11 (7.7%) BC: yes (clinical only) Age: NR Gender (per cent men): NR Severity: PASI = 3.75 (2.90SD) INCLUSION CRITERIA • People with mild to moderate psoriasis EXCLUSION CRITERIA • NR
Interventions	 Calcitriol ointment BD (C) Diflucortolone valerate OM, plus calcitriol ointment ON (C-D)
Outcomes	1. PASI (scale NR)
Notes	Sponsor: NR The trial was set in Korea.
Risk of bias	

Lee 2007 (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	7.7%
Baseline assessments	Unclear risk	Clinical assessments were reported; demo- graphics were unclear
Baseline comparability demonstrated	Unclear risk	Only clinical comparability was demon- strated.

Lepaw 1978

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: predetermined schedule using identical containers coded with participant number Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 29 Treatment duration: 2 wks; FU: 2 wks LF: 2 (6.9%) BC: inadequately reported Age: 14 to 75 Gender (per cent men): 55.2% Severity: not reported INCLUSION CRITERIA • Bilaterally similar psoriatic lesions of the scalp • Adults or adolescents EXCLUSION CRITERIA • Systemic therapy, topical scalp treatments
Interventions	 Halcinonide solution 0.1% TDS (H) Placebo (vehicle) TDS (P)

Lepaw 1978 (Continued)

Outcomes	 Overall therapeutic response Overall comparative response
Notes	The trial did not report sponsorship. This was a scalp trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	6.9%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Levine 2010 (H)

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation table Concealment: adequate BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Double-blind (participant/investigator)
Participants	N: 168 Treatment duration: 12 wks; FU: 12 wks LF: 4 (2.4%) BC: stated, not demonstrated Age: not reported Gender (per cent men): not reported Severity: TLPSS = 6.69 (1.38 SD) INCLUSION CRITERIA • People aged 18 and over with moderate plaque psoriasis

Levine 2010 (H) (Continued)

	 Bilateral plaques or 2 plaques > = 5 cm distant on same side 15 x 15 cm>plaque size > 2 x 2 cm EXCLUSION CRITERIA BSA > 15% Participants with only scalp, nail, flexural, palmoplantar, or pustular psoriasis Participation in trial within previous 3 mths Topical antipsoriatic therapy within previous 2 wks Systemic antipsoriatic therapy within previous 1 mth Use of systemic niacin or multivitamins within previous 2 wks Concomitant use of carbamazepine or primidone People starting or modifying treatment with betablockers within previous 1 mth Significant hematologic, renal, or liver disease Relevant psychiatric or medical illness or surgery Hemoglobin < 10.0 g/dL, hematocrit < 30%, white blood cell count < 3000/mcg/L, platelets < 100,000/mcg/L, aspartate aminotransferase or alanine aminotransferase > 3 x upper limit of normal Pregnancy
Interventions	 Participants were randomised to 2 of 7 treatments: calcipotriene 0.005% ointment BD (C) placebo ointment BD (P)] nicotinamide 1.4% BD (NI) calcipotriene 0.005% ointment BD + nicotinamide 0.05% BD (NC005) calcipotriene 0.005% ointment BD + nicotinamide 0.1% BD (NC010) calcipotriene 0.005% ointment BD + nicotinamide 0.7% BD (NC070) calcipotriene 0.005% ointment BD + nicotinamide 1.4% BD (NC140)
Outcomes	 Total Local Psoriasis Severity Score (0 to 12): sum of erythema, plaque elevation, scaling (each scored 0 to 4) Efficacy end-point: per cent patients 'clear to almost clear' (TLPSS = 0 to 2): 'clear': no elevation above normal skin, no scaling, and no erythema 'almost clear': no elevation above normal skin, surface dryness with some white coloration, and slight to moderate red colouration
Notes	Dermipsor Ltd sponsored the trial. Compliance rates (medicine usage) were reported for each regimen The trial author supplied unpublished data. As participants were randomised to 2 of 7 treatments, some comparisons were within- patient and others between-patient

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	The participants, investigators, study site personnel, sponsor, and laboratories were unaware of the treatment assignments; throughout the study, the study investiga- tor retained a set of code envelopes that

Levine 2010 (H) (Continued)

		were to be opened by the investigator only if deemed necessary after a serious adverse event
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	A computer-generated randomisation table was used.
Loss to follow up	Low risk	2.4%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	Clinical comparability was stated (not demonstrated).

Levine 2010 (P)

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation table Concealment: adequate BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 168 Treatment duration: 12 wks; FU: 12 wks LF: 4 (2.4%) BC: stated, not demonstrated Age: not reported Gender (per cent men): not reported Severity: TLPSS = 6.69 (1.38SD) INCLUSION CRITERIA People aged 18 and over with moderate plaque psoriasis Bilateral plaques or 2 plaques > = 5 cm distant on same side 15 x 15 cm>plaque size > 2 x 2 cm EXCLUSION CRITERIA BSA > 15% Participants with only scalp, nail, flexural, palmoplantar, or pustular psoriasis Participation in trial within previous 3 mths Topical antipsoriatic therapy within previous 1 mth

Levine 2010 (P) (Continued)

	 Use of systemic niacin or multivitamins within previous 2 wks Concomitant use of carbamazepine or primidone People starting or modifying treatment with betablockers within previous 1 mth Significant hematologic, renal, or liver disease Relevant psychiatric or medical illness or surgery Hemoglobin < 10.0 g/dL, hematocrit < 30%, white blood cell count < 3000/mcg/L, platelets < 100,000/mcg/L, aspartate aminotransferase or alanine aminotransferase > 3 x upper limit of normal Pregnancy
Interventions	 Participants were randomised to 2 of 7 treatments: calcipotriene 0.005% ointment BD (C) placebo ointment BD (P)] nicotinamide 1.4% BD (NI) calcipotriene 0.005% ointment BD + nicotinamide 0.05% BD (NC005) calcipotriene 0.005% ointment BD + nicotinamide 0.1% BD (NC010) calcipotriene 0.005% ointment BD + nicotinamide 0.7% BD (NC070) calcipotriene 0.005% ointment BD + nicotinamide 1.4% BD (NC140)
Outcomes	 Total Local Psoriasis Severity Score (0 to 12): sum of erythema, plaque elevation, scaling (each scored 0 to 4) Efficacy end-point: per cent patients 'clear to almost clear' (TLPSS = 0 to 2): 'clear': no elevation above normal skin, no scaling, and no erythema 'almost clear': no elevation above normal skin, surface dryness with some white coloration, and slight to moderate red colouration
Notes	Dermipsor Ltd sponsored the trial. Compliance rates (medicine usage) were reported for each regimen The trial author supplied unpublished data. As participants were randomised to 2 of 7 treatments, some comparisons were within- patient and others between-patient

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	The participants, investigators, study site personnel, sponsor, and laboratories were unaware of the treatment assignments; throughout the study, the study investiga- tor retained a set of code envelopes that were to be opened by the investigator only if deemed necessary after a serious adverse event
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Levine 2010 (P) (Continued)

Randomisation method reported	Low risk	A computer-generated randomisation table was used.
Loss to follow up	Low risk	2.4%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	Clinical comparability was stated (not demonstrated).

Liao 2007

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 50 Treatment duration: 6 wks; FU: 6 wks LF: 1 (2.0%) BC: yes Age: 39.6 (12.8SD); range 21 to 69 Gender (per cent men): 73.5% Target lesion: face (89.8%); genitofemoral (10.2%) Severity: TAS (0 to 12) = 6.2 (3.32SD) INCLUSION CRITERIA People aged 18 to 70 with chronic plaque psoriasis affecting facial and genitofemoral areas EXCLUSION CRITERIA Topical therapies within previous 1 wk UV therapy or sunbathing within previous 2 wks Systemic therapy within previous 4 wks Concurrent use of oral calcium or Vitamin D supplements, or lithium, beta-blockers, or ACE inhibitors
Interventions	 Calcitriol 3 mcg/g ointment BD (C) Tacrolimus 0.3 mg/g ointment BD (T) No concomitant topical therapies or emollients were permitted. Dosing frequency could be reduced or medication discontinued depending on level of irritancy

Liao 2007 (Continued)

Outcomes	 Target Area Score (TAS): 0 to 12 (erythema, plaque elevation, scaling) Physician Global Assessment of improvement (PGA) on a 7-pt scale (worse to clear) Safety: erythema (0 to 4), perilesional oedema (0 to 4), stinging/burning (0 to 4), hot sensation (0 to 4), folliculitis/acne (0 to 1)
Notes	Galderma, Taiwan, sponsored the trial. Individual safety measures were reported, but the total number of participants experi- encing any adverse event were not reported The trial author supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	2.0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Lin 2007

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 14 Treatment duration: 8 wks; FU: 8 wks LF: 4 (28.6%) BC: yes Age: 35.8 (10.4SD), range = 21 to 54

	Gender (per cent men): 78.6%
	Severity: PSI (0 to 12) = 9.9 (1.2SD); PASI = 11.7 (4.8SD), range = 5.5 to 19.2; per
	cent BSA = 13.6 (6.1SD), range = 3% to 20%
	Duration (yrs): 110.8 (9.4SD); range = 2 to 30
	INCLUSION CRITERIA
	• Adult people with recalcitrant, bilateral chronic plaque psoriasis affecting 5% to 20% of BSA
	• Plaques 3 to 20 cm diameter
	• Disease duration > = 2 yrs
	• Resistant to > = 1 topical therapy
	• No other significant medical problem
	EXCLUSION CRITERIA
	• Systemic therapy within previous 30 days
	• Topical therapy within previous 14 days
	• Tests positive for HIV, hepatitis B, or hepatitis C
	• Known sensitivity to Chinese herbs
	• Pregnancy or risk thereof
	• Lactation
	Clinically significant laboratory abnormality
Interventions	• Indigo naturalis 1.4% ointment,OD (I)
	• Placebo ointment (olive oil, yellow wax, petroleum jelly) OD (P)
Outcomes	1. Clearing per cent area of target lesions affected
	2. Psoriasis severity Index (TSS, 0 to 12)
	3. PASI (range not stated)
	4. Safety (immunohistochemical analysis of skin biopsies and blood tests)
Notes	The sponsor was not reported.
	Assessments were conducted by 2 blinded investigators.
	A 3-month uncontrolled follow-study also conducted.
	The trial author supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	High risk	28.6%
Baseline assessments	Unclear risk	These were reported.

Baseline comparability demonstrated	Unclear risk	This was demonstrated.
Lin 2008		
Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described	
Participants	WITHDRAWAL/DROPOUT	
Interventions	 Indigo naturalis 1.4% ointment OD (I) Placebo ointment (olive oil, yellow wax, petroleum jelly) OD (P) Participants achieving clearance before study end point could stop treatment 	

Lin 2008 (Continued)

Outcomes	 Global improvement (IAGI); 6-pt: worse to clearance Signs (erythema, induration, scaling), each scored 0 (none) to 8 (very severe) Total sum score (0 to 24) PASI (range unclear): baseline median used to explore response by group Compliance (medication use) Safety
Notes	The trial was supported by a grant: CMRPG33024 from Chang Gung Memorial Hos- pital Participants were instructed to wash skin thoroughly before visits; assessments were based on photographs

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Unclear risk	These were reported.
Baseline comparability demonstrated	Unclear risk	This was demonstrated.

Lister 1997

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not stated
	Concealment: unclear
	BLINDING
	Single-blind (investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 171
1 articipants	Treatment duration: 8 wks; FU: 16 wks
	LF: not reported
	BC: psoriasis comparable, demographics unclear
bC. psonasis comparable, demographics unclear	

Lister 1997 (Continued)

Interventions	Age: not stated Gender (per cent men): not stated Severity: TSS (scale NR) = 6.24 INCLUSION CRITERIA • Chronic plaque psoriasis EXCLUSION CRITERIA • Unclear • Dithranol cream 1 to 3% OD (D)
Outcomes	 Calcipotriol BD (C) 1. Total Sign Score (erythema, scaling, induration) (scale NR) 2. Investigator and Patient Global Assessments (scales NR) 3. Relapse rates
Notes	Bioglan Laboratories sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient detail.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Lowe 2005

Methods DESIGN Between-patient Participant delivery ALLOCATION Random Random Method of randomisation: consecutive balanced blocks of 7 (3: Concealment: unclear BLINDING Single-blind (investigator); as cream and lotion were compared patients WITHDRAWAL/DROPOUT Described Participants N: 192 Treatment duration: 4 wks; FU: 8 wks LF: 0 (0%) BC; yes Age: 48.65; range = 19 to 79 Gender (per cent men): 65.6% Ethnicity (per cent white): 82.8% Severity: DSS (to 12) 2 = 7.60 (1.55D) INCLUSION CRITERIA • People aged > = 18 with stable moderate to severe plaque p • DSS (to 12) 2 = 66 • Target lesion diameter > 3 cm, lesion not located on face, a hands, or feet EXCLUSION CRITERIA • Clobetasol propionate lotion/cream BD (CP) • Placebo lotion BD (P) Participants were randomised to clobetasol propionate cream aggregated for review purposes Other affected areas could also be treated. There was a 4-wk treatment- free follow-up period. Outcomes 1. Signs: erythema, plaque elevation, scaling, prurius (0 (non 2. Dermatological sum score (DSS) (crythema, plaque elevati 3. Investigator's Assessment of Global improvement (LGI): 7. • Clobeta severity score (GSS): dichotomised as success (GSS (GSS > 2)) <th>DESIGN</th> <th>г</th> <th></th>	DESIGN	г	
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Method of randomisation: consecutive balanced blocks of 7 (3:: Concealment: unclear BLINDING Single-blind (investigator); as cream and lotion were compared patients WTTHDRAWAL/DROPOUT Described Participants N: 192 Treatment duration: 4 wks; FU: 8 wks LF: 0 (0%) BC: yes Age: 48.65; range = 19 to 79 Gender (per cent men): 65.6% Ethnicity (per cent white): 82.8% Severity: DSS (0 to 12) = 7.60 (1.55SD) INCLUSION CRITERIA • People aged > = 18 with stable moderate to severe plaque p • DSS (0 to 12) = 5 6 • Target lesion diameter > 3 cm, lesion not located on face, a hands, or feet EXCUSION CRITERIA • Cobetasol propionate lotion/cream BD (CP) • Placebo lotion BD (P) Participants were randomised to clobetasol propionate cream aggregated for review purposes Other affected areas could also be treated. There was a 4-wk treatment- free follow-up period. Outcomes 1. Signs: erythema, plaque elevation, scaling, pruritus (0 (nor 2. Dermatological sum score (DSS) (crythema, plaque elevatio 3. Investigator's Assessment of Global improvement (LGGI); 7 • Global severity score (GSS) (ci			
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Single-blind (investigator); as cream and lotion were compared patients WITHDRAWAL/DROPOUT Described Participants N: 192 Treatment duration: 4 wks; FU: 8 wks LF: 0 (0%) BC: yes Age: 48.65; range = 19 to 79 Gender (per cent men): 65.6% Ethnicity (per cent white): 82.8% Severity: DSS (0 to 12) = 7.60 (1.55SD) INCLUSION CRITERIA • People aged > = 18 with stable moderate to severe plaque p • DSS (to 12) >= 6 • Target lesion diameter > 3 cm, lesion not located on face, a hands, or feet EXCLUSION CRITERIA • Cobstasol propionate lotion/cream BD (CP) • Placebo lotion BD (P) Participants were randomised to clobetasol propionate cream aggregated for review purposes Other affected areas could also be treated. There was a 4-wk treatment- free follow-up period. Outcomes 1. Signs: erythema, plaque elevation, scaling, pruritus (0 (non 2. Dermatological sum score (DSS) (erythema, plaque elevati a. Investigator's Assessment of Global improvement (LAGI); 7. 4. Global severity score (GSS): dichotomised as success (GSS (GSS > = 2) 5. BSA Salderma R&D, Sophia Antipolis, France, sponsored the trial.			
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Risk of bias			
Bias Authors' judgement Support for judge	Authors' judgem	A	Support for judgement

Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator); as the cream and lotion were compared, it was not possible to blind participants
Randomisation method reported	Low risk	Participants allocated in consecutive bal- anced blocks of 7 using the following ra- tios: 3:3:1
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Luger 2008

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not stated
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Danticipants	N: 869
Participants	Treatment duration: 52 wks; FU: 52 wks
	LF: 55 (6.3%)
	BC: yes
	Age: $48.7 (15.0$ SD), range = $18 \text{ to } 86$
	Gender (per cent men): 44.0%
	Ethnicity (per cent white): 96.7%
	Severity: IGA moderate = 55.5%; IGA severe = 37.8%; IGA very severe = 6.7%
	Duration (yrs): 17.6 (14.2SD)
	INCLUSION CRITERIA
	• Hospital outpatients aged > = 18 with moderate to severe scalp psoriasis
	 Amendable to topical treatment with <= 100 g/wk
	Psoriasis vulgaris on trunk or limbs;
	 Area of scalp affected > 10%
	 IGA at least moderate severity
	EXCLUSION CRITERIA
	• Use within previous 28 days of PUVA or grenz ray therapy
	• Systemics with a potential impact on scalp psoriasis, or biological therapies; UVB;

	topical therapies; calcium metabolism disorders associated with hypercalcaemia. Unclear if concurrent use permitted during the trial
Interventions	 Calcipotriol 50 mcg/g in scalp formulation (gel) OD PRN (C) Calcipotriol gel 50 mcg/g plus betamethasone dipropionate "scalp formulation" (gel) 0.5 mg/g OD (C-B) Participants achieving clearance could stop treatment, but remained in the trial, and restart as needed. The regimen was intended to reflect usual clinical practice
Outcomes	 Adverse drug reactions Adverse events associated with long-term corticosteroid use Investigator's Global Assessment of disease severity (IGA) (6-pt: absence of disease to very severe disease (5)) Controlled disease: IGA < = 2 Compliance (self-report; failure to use treatment for any reason other than no treatment required) (tube weight also assessed)
Notes	Leo Pharma, Ballerup, Denmark, sponsored the trial. Adverse events were recorded by investigators and reviewed by an adjudication committee (Data Safety Monitoring Board) comprising of 3 dermatologists blinded to treatment assignment The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	This was not stated.
Loss to follow up	Low risk	6.3%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Maier 2004

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT
	Not described
Participants	N: 34 Treatment duration: 8 wks; FU: 8 wks LF: 10 (29.4%) BC: yes (clinical only) Age: NR Gender (per cent men): 55.9% Severity: mPASI = 6.3 (2.2SD) INCLUSION CRITERIA • People with stable mild to moderate plaque psoriasis EXCLUSION CRITERIA • Concomitant antipsoriatic therapy or use within previous 2 wks
Interventions	 Herbal skincare products (Dr Michaels® cleansing gel, ointment, and skin conditioner) BD (H) 3 fatty skin care products (not vehicle) BD (P)
Outcomes	1. mPASI (0 to 64.8)
Notes	The trial did not state the sponsor. This was an abstract only.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	High risk	29.4%
Baseline assessments	Unclear risk	Clinical assessments were reported.

Baseline comparability demonstrated Unclear risk Clinical comparability was demonstrated. Medansky 1987 DESIGN Methods Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described Participants N: 121 Treatment duration: 3 wks; FU: 3 wks LF: 6 (5.0%) BC: yes, except duration of disease (P = 0.04) Age: 53.8 (range = 16 to 80) Gender (per cent men): 67.8% Severity: TSS (0 to 9) = 6.6 Duration (yrs): 17.9 (range = 1 to 52) **INCLUSION CRITERIA** • Aged \geq 12; chronic plaque psoriasis, stable, or worsening • Duration ≥ 1 year • Total Sign Score ≥ 6 **EXCLUSION CRITERIA** • Concomitant medication • Recent systemic corticosteroids or antimetabolites • Recent topical corticosteroids • Pregnancy • Lactation • Those needing > 90 g/wk topical steroid Other forms of psoriasis Interventions • Mometasone furoate ointment 0.1% OD (M) • Vehicle OD (P) Outcomes 1. Signs (erythema; induration; scaling) 2. Total Sign Score (0 to 9) 3. Investigator Global Assessment (6-pt: no change or worse to cleared or marked improvement) Notes The trial was supported in part by a grant from Schering Corporation. There was SD imputation (TSS).

Topical treatments for chronic plaque psoriasis (Review)

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Medansky 1987 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	67.8%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	Baseline comparability was demonstrated (only significant difference: duration of dis- ease)

Medansky 1996

Methods	DESIGN
	Within-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: "schedule"
	Concealment: unclear
	BLINDING
	Double-blind (participant/assessor)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 134
F	Treatment duration: 3 wks; FU: 3 wks
	LF: 6 (4.5%)
	BC: unclear
	Age: 47 (range = 20 to 81)
	Gender (per cent men): not stated
	Severity: TSS (0 to 9) = 6.5; per cent unstable psoriasis = 28%
	INCLUSION CRITERIA
	• Mild-to-moderate symmetrical chronic plaque psoriasis
	• Adult
	• TSS (0 to 9) ≥ 6
	EXCLUSION CRITERIA
	• Recent topical or systemic antipsoriatic therapy
	• Recent lithium, NSAIDs, or beta-blockers

Medansky 1996 (Continued)

Interventions	 Diflorasone diacetate ointment, 0.05% BD (D) Calcipotriene ointment 0.005% BD (C)
Outcomes	 Signs (erythema, scaling, induration) Total Sign Score (0 to 9) Physician overall evaluation (7 pt: worse to clear) Physician comparative evaluation Patient comparative evaluation
Notes	Dermik Laboratories Inc. sponsored the trial. Adverse events: itching and burning There was SD imputation (TSS/IAGI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	4.5%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Menter 2009

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Open (blinding NS but product vehicles differ)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 122 Treatment duration: 4 wks; FU: 8 wks LF: 0 (0%)

Menter 2009 (Continued)

	BC: yes Age: 46.5 (14.2SD) Gender (per cent men): 61.1% Duration (yrs): 16.4 (11.50SD) Severity: ODS = 3.06 (0.43SD) (participant demographics based on per-protocol sample). INCLUSION CRITERIA • People aged 18 to 80 with moderate to severe stable plaque psoriasis • BSA: 3 to 20% EXCLUSION CRITERIA • Known allergy to study ingredients • Psoriasis affecting scalp, face, or groin only • Area affected required weekly treatment with > 50 g CP spray or > 100 g C-BD
	• Area affected required weekly treatment with > 50 g Cr spray of > 100 g C-BD ointment
Interventions	 Calcipotriol 50 mcg/g + betamethasone dipropionate 0.5 mg/g combination ointment OD (C-BD) Clobetasol propionate 0.05% spray (CP)
Outcomes	 Overall disease severity (ODS) on a 5-pt scale: clear (0), almost clear, mild, moderate, severe/very severe (4) Based on erythema, scaling, and elevation Investigator Global Assessment (IGA) on a 4-pt scale (0 to 3): clear (0), mild, moderate, severe (3). Based on per cent BSA affected (extent) rather than signs
Notes	Galderma Laboratories, LP, sponsored the trial. Atrophy was not assessed. Telangiectasia was not assessed. 29 participants were 'non-compliant' with treatment, but intention-to-treat (ITT) anal- ysis included all enrolled participants The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open (blinding not reported, but product vehicles differ)
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.

Menter 2009 (Continued)

Baseline comparability demonstrated	Low risk	This was demonstrated.
Molin 1997		
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not Concealment: unclear BLINDING Double-blind (participant/asses WITHDRAWAL/DROPOUT Described	ssor)
Participants	N: 421 Treatment duration: 8 wks; FU LF: 4 (1%) BC: Psoriasis comparable, demo Age: not stated Gender (per cent men): not stat Severity: no summary measure INCLUSION CRITERIA • Adult outpatients • Mild-to-moderate stable a EXCLUSION CRITERIA • None reported	ographics not reported ted
Interventions	Calcipotriol cream 50 mcgBetamethasone 17-valerate	
Outcomes	•	ment of response (5-pt: worse to cleared) s of response (5-pt: worse to cleared)
Notes	Leo Pharmaceutical Products sp	ponsored the trial

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).

Molin 1997 (Continued)

Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	1.0%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Monastirli 2000

Methods	DESIGN Between-patient Delivery unclear ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Described
Participants	N: 70 Treatment duration: 10 wks; FU: 10 wks LF: not reported BC: yes Age: 45.7 Gender (per cent men): 57.1% Severity: PASI = • D: 7.31 (1.79SD, N = 35) • C: 6.29 (1.63SD, N = 35) Duration (yrs): 17 INCLUSION CRITERIA • Inpatients with chronic plaque psoriasis EXCLUSION CRITERIA • Pregnancy • Lactation • Ineffective contraception • Systemic treatment within previous 2 mths • Hepatic or renal disease • Hypercalcaemia • Known hypersensitivity to study medications
Interventions	 Dithranol 2% 30 minutes OD (D) Calcipotriol ointment 50 mcg/g BD (C)
Outcomes	1. PASI (excluding head)

Monastirli 2000 (Continued)

Notes	The trial did not report sponsorship. Treatment was delivered in an inpatient set	ting.
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias)	High risk	The trial was open.

All outcomes		
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Mortensen 1993b

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
D · · ·	
Participants	N: 34
	Treatment duration: 3 wks; FU: 4 wks
	LF: 0 (0%)
	BC: psoriasis comparable, demographics inadequately reported
	Age: 43 (range = 26 to 75)
	Gender (per cent men): 58.8%
	Severity: PASI = 12.2
	INCLUSION CRITERIA
	• Stable plaque-type psoriasis
	Adult outpatients
	Normal hepatic and renal function
	EXCLUSION CRITERIA
	• Recent UV or other psoriasis treatments

Mortensen 1993b (Continued)

	• Disease or medication influencing calcium or bone metabolism
Interventions	 Calcipotriol ointment 50 mcg/g BD (C) [max 100 g/wk] Placebo (vehicle) (P)
Outcomes	 PASI Investigator Global Assessment: per cent improvement from baseline, based on PASI Patient Global Assessment : per cent improvement from baseline, based on PASI
Notes	Leo Pharmaceuticals, Denmark Duration (yrs) sponsored the trial

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Olsen 1991

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 378 Treatment duration: 2 wks; FU: 3 wks LF: 1 (0.3%) BC: yes

Olsen 1991 (Continued)

	Age: 46 (range = 18 to 88) Gender (per cent men): 45% Severity: 80% moderate ($6 \le TSS < 7.5$), 20% severe ($TSS > 7.5$) Duration (yr): 12.0 (9.7SD, N = 377); range = 0.4 to 55.0 INCLUSION CRITERIA • Moderate to severe scalp psoriasis (TSS (0 to 9) \ge 6) • Stable or worsening • Adult EXCLUSION CRITERIA • Recent systemic, topical, or UV treatment for psoriasis.
Interventions	 Clobetasol propionate 0.05% BD (C) Placebo (vehicle) (P)
Outcomes	 Severity (erythema; induration; scaling, pruritis) TSS (0 to12) Investigator Global Assessment (6-pt: worse to cleared) Patient Global Assessment (4-pt: poor to excellent)
Notes	In part, Glaxo Inc. sponsored the trial. This was a scalp trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.3%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Olsen 1996 (1)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described (for both trials together)
Participants	N: 181 Treatment duration: 4 wks; FU: 4 wks LF: 3 (1.7%) BC: yes Age: 49 (range = 15 to 76) Gender (per cent men): 68.0% Severity: per cent BSA affected = 12.0% (range = 1% to 80%); per cent BSA treated = 11% (range = 1% to 80%) Duration (yrs): 19 (range = 1 to 60)
Interventions	 Fluticasone propionate 0.005% ointment (F) (max 100 g/wk) Placebo (vehicle) (P)
Outcomes	 Investigator Global Assessment (6-pt: cleared to worse) Severity (erythema; induration; scaling; pruritis) Patient subjective assessment (treatment effect: 1 = excellent to 4 = poor)
Notes	The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	1.7%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Olsen 1996 (2)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described (for both trials together)
Participants	N: 207 Treatment duration: 4 wks; FU: 4 wks LF: 2 (1.0%) BC: yes Age: 45 (range = 12 to 87) Gender (per cent men): 52.7% Severity: per cent BSA affected = 12.5% (range = 1% to 80%); per cent BSA treated = 12% (range = 1% to 80%) Duration (yrs): 16 (range = 0.8 to 52)
Interventions	 Fluticasone propionate 0.005% ointment (F) (max 100 g/wk) Placebo (vehicle) (P)
Outcomes	 Investigator Global Assessment (6-pt: cleared to worse) Severity (erythema; induration; scaling; pruritis) Patient subjective assessment (treatment effect: 1 = excellent to 4 = poor)
Notes	The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	1.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Oranje 1997

Methods Participants	DESIGN Between-patient Patient/parent delivery ALLOCATION Random Method of randomisation: computer-generated random number table used by 7/14 centres to randomly select ≤ 3 participants Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described N: 77 Treatment duration: 8 wks; FU: 8 wks LF: 0 (0%)	
	 LF: 0 (0%) BC: yes Age: 10 (range = 2 to 14) Gender (per cent men): 46.8% Severity: not reported INCLUSION CRITERIA Mild to moderate chronic plaque psoriasis Children aged 2 to 14 EXCLUSION CRITERIA Acute guttate, pustular, erythrodermic, or worsening psoriasis Psoriasis mainly on the face, scalp, or diaper area Systemic treatment Recent phototherapy Concurrent Vitamin D, calcium, or other intercurrent medication Renal, hepatic, or osteoarthritic disease 	
Interventions	 Calcipotriol ointment 50 mcg/g BD (C) Placebo (vehicle) (P) 	
Outcomes	 PASI: severity (redness; thickness; scaliness, area) Extent of disease Investigator Global Assessment Patient Global Assessment (by parent/guardian for those aged < 8) Compliance 	
Notes	Leo Pharmaceutical Products, Denmark, sponsored the trial. The trial participants were children.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Oranje 1997 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Ormerod 1997

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Not described
Participants	N: 12 Treatment duration: 2 wks; FU: 2 wks LF: unclear BC: unclear Age: not reported Gender (per cent men): not reported Severity: TSS (0 to 24) =12.2 INCLUSION CRITERIA • Bilaterally similar chronic • Stable plaque psoriasis. EXCLUSION CRITERIA • Recent systemic or UV therapy
Interventions	 Betamethasone valerate ointment 0.1% BD (B) White soft paraffin BD (P)
Outcomes	 Signs (erythema; elevation; scaling) Total Sign Score (0 to 24)
Notes	Wyeth-Ayerst Research and Glaxo Dermatology sponsored the trial

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Ormerod 2000

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING	
Participante	Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 17 Treatment duration: 4 wks; FU: 4 wks LF: 0 (0%) BC: not reported Age: not reported Gender (per cent men): not reported Severity: TSS (0 to 24) = 15.5 (SD: 3.7; N = 17) INCLUSION CRITERIA • Stable plaque psoriasis EXCLUSION CRITERIA • Pregnancy • Ineffective contraception • Lactation	

Ormerod 2000 (Continued)

Interventions	 NG-monomethyl-L-arginine (L-NMMA) cream 25% BD NG-monomethyl-L-arginine (L-NMMA) cream 5% BD Placebo (P) 	
Outcomes	1. TSS (elevation, erythema, scaling) (0 to 24)	
Notes	The trial did not report sponsorship.	

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Ormerod 2005

Methods	DESIGN Within-patient Participant delivery ALLOCATION	
	Random Method of randomisation: randomised in blocks of 4 by the pharmacy department using	
	'Minitab'	
	Concealment: unclear	
	BLINDING	
	Double-blind (participant/investigator)	
	WITHDRAWAL/DROPOUT	
	Unclear	
Participants	N: 24	
	Treatment duration: 12 wks; FU: 12 wks	
	LF: 2 (8.3%)	
	BC: yes	
	Age: 47.5 (range = 28 to 78)	
	Gender (per cent men): 75.0%	
	Severity: TSS (0 to 24) = 17.0 (3.3SD, N = 22)	

Ormerod 2005 (Continued)

	INCLUSION CRITERIA
	Chronic plaque psoriasis
	• Aged 18 and over
	• No significant co morbidity
	• Transminase levels within double normal upper limit
	EXCLUSION CRITERIA
	• Planned exposure to sunlight over trial duration
	• Pregnancy or risk thereof
	• Lactation; systemic or UV therapy within previous 4 wks
	• Topical therapy within previous 2 wks
	 Known allergy to macrolide drugs
	Renal or hepatic disease
	• Renal malignancy within previous 5 yrs
Interventions	 Topical sirolimus 2.2% for 6 wks then 8% for a further 6 wks (S) Placebo (P) Dosing frequency was not reported.
Outcomes	1. Total Sign Score (TSS) of target plaque (erythema, thickening, scaling) (0 to 24)
Notes	Wyeth Research, Philadelphia, sponsored the trial. No systemic adverse event was considered clinically significant. The daily dosing frequency was unclear.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Block randomisation was used.
Loss to follow up	Low risk	9.3%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Ortonne 1994

Methods	DESIGN Between-patient	
	Participant delivery	
	ALLOCATION Random	
	Method of randomisation: participants allo	ocated sequentially upon inclusion
	Concealment: unclear	searced sequencially upon inclusion
	BLINDING	
	Double-blind (participant/assessor)	
	WITHDRAWAL/DROPOUT	
	Described	
Participants	N: 188	
	Treatment duration: 6 wks; FU: 6 wks	
	LF: 32 (17.0%)	
	BC: yes	
	Age: 46.0 (range = 20 to 85)	
	Gender (per cent men) 67.3%	(7
	Severity: per cent BSA = 18.9%; PASI = 11.67; per cent unstable psoriasis = 40% Duration (yrs): 14.9 INCLUSION CRITERIA • Chronic plaque psoriasis	
	Stable or worsening	
	• BSA 10% to 40%	
	• PASI 1 to 30; outpatients	
	EXCLUSION CRITERIA	
	 Pregnancy Lactation	
	 Lactation Concurrent disease	
	Concomitant therapy	
	• Hypersensitivity to Vitamin D or ana	logues
	• Planned exposure to sun	
Interventions	 Calcipotriol ointment BD (C) Calcipotriol ointment OM, plus betamethasone dipropionate ointment ON (CB) 	
2		
Outcomes	 PASI Investigator Global Assessment 	
	2. Investigator Global Assessment	
Notes	Leo Pharmaceuticals sponsored the trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Ortonne 1994 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Low risk	Randomisation was sequential.
Loss to follow up	Low risk	17.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Ortonne 2003

Methods	 DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation by blocks of 4 using RANUNI routine of the SAS system Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 75 Treatment duration: 6 wks; FU: 6 wks LF: 10 (13.3%) BC: yes, within-patient design but comparability of lesions at baseline not reported Age: 44.5 (14.5SD: range = 18.8 to 70.7) Gender (per cent men): 53% Severity: not reported INCLUSION CRITERIA People with stable chronic plaque psoriasis, localised on 'sensitive areas': face, hairline, retro-auricular, and flexural areas Aged 18 to 70 1 to 4 bilateral lesions of similar severity EXCLUSION CRITERIA Pregnancy or risk thereof Lactation Concomitant disease Acute guttate, pustular, erythrodermic, or arthropathic psoriasis History of hypercalcaemia, renal dysfunction Calcium-based calculi Conditions requiring systemic supplements of vitamin D or calcium Previous topical therapy within previous 2 wks (4 wks for retinoids)

Ortonne 2003 (Continued)

	• Previous systemic therapy within previous 4 wks (16 wks for retinoids)	
Interventions	 Calcitriol ointment 3 mcg/g BD (C1) Calcipotriol ointment 50 mcg/g BD (C2) 	
Outcomes	 Investigator's Global Assessment of local safety for each lesion (3-pt: 0 = poor to 2 = excellent) Mean of worst sign scores (0 to 9) (signs: perilesional erythema; perilesional oedema; stinging/burning) Investigator's Global Assessment of Improvement (IAGI) (7-pt: worse to clear) Patient's preference for tolerance, efficacy, and global preference (5-pt: right-hand side (RHS) > left-hand side (LHS): -2 to LHS > RHS: 2) 	
Notes	The trial did not report sponsorship, but 2 authors worked for Galderma, France This was an inverse psoriasis trial. There was SD imputation (TSS).	

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was single-blind (investigator).
Randomisation method reported	Low risk	A computer-generated block list was used for randomisation.
Loss to follow up	Low risk	13.3%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Ortonne 2004

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation schedule Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 501 Treatment duration: 8 wks; FU: 8 wks LF: 21 (4.2%) BC: yes Age: 51.2 (15.0SD, N = 501) Gender (per cent men): 54.9% Severity: PASI = 9.8 (6.1SD, N = 501) Duration (yrs): 19.4 (14.6SD, N = 501) INCLUSION CRITERIA • Stable chronic plaque psoriasis amenable to topical treatment • Aged 18 and over EXCLUSION CRITERIA • Pregnancy or risk thereof • Lactation • Unstable psoriasis or other inflammatory diseases • Abnormality of calcium metabolism or hypercalcaemia • Systemic or phototherapy within previous 4 wks • Topical therapy within previous 2 wks • Other topical therapy for trunk or limbs during study period • Corticosteroid treatment of scalp (WHO: class IV) or facial area (WHO: class III/ IV) during study period	
Interventions	 TCP ointment ON for 4 wks then calcipotriol ointment 50 mcg/g ON for 4 wks (A) Tacalcitol ointment 4 mcg/g ON for 8 wks (T) TCP: calcipotriol 50 mcg/g, plus betamethasone dipropionate 0.5 mg/g ointment 	
Outcomes	 PASI: mean % reduction IAGI (6-pt: worse to clearance) PAGI (6-pt: worse to clearance) 	
Notes	Leo Pharmaceutical Products sponsored the trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement

Ortonne 2004 (Continued)

Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	4.2%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Ortonne 2006

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: generated by sponsor; stratified by centre (N = 16); Concealment: unclear BLINDING Single-blind (investigator); as cream and gel were compared, it was not possible to blind participants WITHDRAWAL/DROPOUT
Participants	DescribedN: 124Treatment duration: 12 wks; FU: 14 wksLF: 0 (0%)BC: yesAge: 48.2 (14.8SD); range = 18 to 82Gender (per cent men): 68.5%Ethnicity (per cent white): 96.0%Severity: per cent BSA = 6.3 ((2.2SD), range = 1% to 10%; LPSI (0 to 12) = 7.2 (1.7SD), range = 4 to 12; PASI (range NS) = 7.1 (3.7SD), range = 1.4 to 21.8Duration (yrs): 18.9 (13.9SD); range = 0.7 to 69.3INCLUSION CRITERIA• People aged > = 18 with mild to moderate stable plaque psoriasis• Per cent BSA < = 10%

Ortonne 2006 (Continued)

	to alter tacrolimus concentrations, lithium, beta blockers, antimalarials, other medicated topical agents
Interventions	 Calcipotriol 0.005% ointment BD (C) Tacrolimus 0.3% gel BD or 0.5% tacrolimus cream BD (T) There were 10 to 14 hours between applications; lesions were treated for 7 days following clearance Participants were randomised to tacrolimus cream or gel; results were aggregated for review purposes
Outcomes	 Local Psoriasis Severity Index (LPSI) (erythema, induration, scaling): 0 to 12 Physician assessment of clinical response of target lesion: IAGI: 5-pt = 0 (worse) to 4 (much better) Subject's assessment of clinical response of target lesion: PAGI: 5-pt = 0 (worse) to 4 (much better) Compliance (median daily medication usage) Cosmetic acceptability (participant) Adverse events and clinical laboratory evaluations
Notes	Fujisawa GmbH, Munich, Germany, sponsored the trial. Vissers 2008 (secondary reference under Ortonne 2006) reports immunohistochemistry findings for subgroup (N = 18)

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator); as the cream and gel were compared, it was not possible to blind participants
Randomisation method reported	Low risk	Randomisation was generated by the spon- sor and stratified by centre
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Ortonne 2010

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: pre-planned computer-generated schedule Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 408 Treatment duration: 8 wks; FU: 8 wks LF: 9 (2.2%) BC: yes Age: 45.6; range = 18 to 86 Gender (per cent men): 54.7% Ethnicity (per cent white): 97.8% Severity: PASI = 9.6, range = 0.9 to 32.7; IGA moderate = 68.6% Duration (yrs): 17.7; range = 0 to 70 INCLUSION CRITERIA People aged > = 18 with chronic plaque psoriasis amenable to topical treatment % face affected >10% % 1 of more body areas affected > = 10% i GA at least mild EXCLUSION CRITERIA Pregnancy or risk thereof Lactation Unstable forms of psoriasis in treatment areas Other skin condition that could affect assessment PUVA or systemic therapy within previous 4 wks UVB or topical therapy on trunk, limbs, or face within previous 2 wks Treatment of scalp psoriasis with very potent (WHO group IV) corticosteroids or vitamin D analogues Known or suspected abnormality in calcium homeostasis
Interventions	 Calcipotriol 25 mcg/g ointment OD (C25) Calcipotriol 25 mcg/g ointment plus hydrocortisone 10 mg/g ointment, OD (C25H) Calcipotriol 50 mcg/g ointment OD (C50) Calcipotriol 50 mcg/g ointment plus hydrocortisone 10 mg/g ointment OD (C50H)
Outcomes	Outcomes were assessed for the face and body overall and for the face only: 1. PASI (per cent change from baseline to EOT): face = 0 to 3.6; face + body = 0 to 68.4 [1] 2. Investigator's Global Assessment of Disease Severity (IGA): 6-pt = absence of disease to very severe disease 3. Adverse events

Ortonne 2010 (Continued)

	4. Compliance (self-report)
Notes	Leo Pharma A/S sponsored the trial. The sponsor supplied unpublished data.

Risk of bias

-		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	A preplanned computer-generated sched- ule was used.
Loss to follow up	Low risk	2.2%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Papakostantinou 2005

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 22 Treatment duration: 2 wks; FU: 2 wks LF: 0 (0%) BC: no Age: 41.3 (3.94SD) Gender (per cent men): 45.5% Severity: NS INCLUSION CRITERIA • People with mild or moderate plaque psoriasis • PASI (0 to 72) < = 10

Papakostantinou 2005 (Continued)

	 Elbow lesions for treatment > = 5 cm diameter EXCLUSION CRITERIA Antipsoriatic therapy within previous 4 wks 	
Interventions	 Theophylline 1% ointment BD (T) Vehicle ointment (20% transcutol, 10% oleic acid) BD (P) 	
Outcomes	1. PASI (0 to 72)	
Notes	The sponsor was not stated.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details (ab- stract only).
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Unclear risk	The trial did not report this (but this was a within-patient study, so demographics will be comparable by definition)
Baseline comparability demonstrated	Unclear risk	This was not demonstrated (see comment above).

Papp 2003 (H)

Methods	
	DESIGN
	Between-patient
	Participant delivery ALLOCATION
	Random
	Method of randomisation: computer-generated random code
	Concealment: adequate
	BLINDING
	Double-blind (participant/assessor) WITHDRAWAL/DROPOUT
	Described
Participants	N: 1043
	Treatment duration: 4 wks; FU: 4 wks
	LF: 15 (1.4%)
	BC: yes
	Age: 47.1
	Gender (per cent men): 58.4%
	Severity: mean PASI = 10.8 (range = 1 to 36)
	Duration: 18.7 years
	INCLUSION CRITERIA
	Chronic plaque psoriasis
	• Aged at least 18
	• BSA $\geq 10\%$
	EXCLUSION CRITERIA
	• Other types of psoriasis or skin diseases
	• Hypercalcaemia
	• Systemic antipsoriatic treatment or UV therapy within previous 6 wks
	• Topical antipsoriatic therapy within previous 2 wks
	• Other concomitant medication that might affect psoriasis
	Contraindications for corticosteroid treatment
	Planned exposure to UV light
	• Pregnancy
	Lactation
Interventions	• Calcipotriol 50 mcg/g + betamethasone dipropionate 0.5 mg/g combination
	ointment BD (D)
	• Calcipotriol 50 mcg/g in combination vehicle ointment BD (C)
	• Betamethasone dipropionate 0.5 mg/g in combination vehicle ointment BD (B)
	• Placebo (combination vehicle) ointment BD (P)
Outcomes	1. PASI (head excluded)
	2. Total Severity Score (9-pt, absent to very severe)
	 IAGI (6-pt: worse to clearance)
	4. PAGI (6-pt: worse to clearance)
Notes	Leo Pharmaceuticals sponsored the trial.

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	Treatments were assigned by computer- generated code and assigned chronologi- cally at each centre
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/ investigator). Treatments were identifiable only by a code number
Randomisation method reported	Low risk	Randomisation was computer-generated. (3:3:3:1)
Loss to follow up	Low risk	1.4%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Papp 2003 (P)

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated random code
	Concealment: adequate
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 1043
T articipants	Treatment duration: 4 wks; FU: 4 wks
	LF: 15 (1.4%)
	BC: yes
	Age: 47.1
	Gender (per cent men): 58.4%
	Severity: mean PASI = 10.8 (range = 1 to 36)
	Duration: 18.7 years
	INCLUSION CRITERIA
	Chronic plaque psoriasis
	 Aged at least 18
	 BSA >10%
	EXCLUSION CRITERIA
	Other types of psoriasis or skin diseases
	 Hypercalcaemia
	• Typercalculation

Papp 2003 (P) (Continued)

	 Systemic antipsoriatic treatment or UV therapy within previous 6 wks Topical antipsoriatic therapy within previous 2 wks Other concomitant medication that might affect psoriasis Contraindications for corticosteroid treatment Planned exposure to UV light Pregnancy Lactation
Interventions	 Calcipotriol 50 mcg/g + betamethasone dipropionate 0.5 mg/g combination ointment BD (D) Calcipotriol 50 mcg/g in combination vehicle ointment BD (C) Betamethasone dipropionate 0.5 mg/g in combination vehicle ointment BD (B) Placebo (combination vehicle) ointment BD (P)
Outcomes	 PASI (head excluded) Total Severity Score (9-pt, absent to very severe) IAGI (6-pt: worse to clearance) PAGI (6-pt: worse to clearance)
Notes	Leo Pharmaceuticals sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	Treatments were assigned by computer- generated code and assigned chronologi- cally at each centre
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator), and treatments were identifi- able only by a code number
Randomisation method reported	Low risk	Randomisation was computer-generated. (3:3:3:1)
Loss to follow up	Low risk	1.4%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Pariser 1996

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Not described
Participants	N: 235 Treatment duration: 8 wks; FU: 8 wks LF: Unclear BC: psoriasis comparable, demographics not reported. Age: 45.1 (range = 18 to 86) Gender (per cent men): not reported Severity: TSS (0 to 9) = 4.75 INCLUSION CRITERIA • Stable plaque-type psoriasis; otherwise healthy, non-pregnant people • At least 4/9 for plaque elevation • BSA range = 5% to 20% EXCLUSION CRITERIA • None reported
Interventions	 Calcipotriene ointment 0.005% OD (C) Placebo (vehicle) (P)
Outcomes	 Severity (scaling; erythema; plaque elevation) TSS (0 to 9) Investigator Global Assessment (10-pt)
Notes	Bristol-Myers Squibb Pharmaceutical Research Institute sponsored the trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.

Pariser 1996 (Continued)

Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Pauporte 2004

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 89 Treatment duration: 3 wks; FU: 4 wks LF: 4 (4.5%) BC: yes Age: 46 (15SD) Gender (per cent men): 44.8% Severity: TSS (0 to 9) = 7.15 INCLUSION CRITERIA • Moderate to severe scalp psoriasis, stable, or slowly exacerbating > 1 wk • Aged ≥ 12 • Good general health • Scalp involvement $\geq 20\%$ • TSS (0 to 9) ≥ 6 EXCLUSION CRITERIA • Pregnancy or risk thereof • Lactation • People requiring topical or systemic treatments that could affect psoriasis • Systemic corticosteroids within previous 4 wks • Topical therapies within previous 1 wk • Concomitant use of other scalp therapies
Interventions	 Fluocinolone acetonide 0.01% topical oil (Derma-Smoothe/FS), plus occlusion ON or for at least 4 hours (F) Placebo oil, plus occlusion ON or for at least 4 hours (P) Participants washed their hair with a non-medicated shampoo after treatment
Outcomes	 Total Severity Score (0 to 9) (erythema, thickening, scaling) Investigator's Assessment of Global Improvement (IAGI) (7-pt: cleared to exacerbation)

Pauporte 2004 (Continued)

Notes	The trial did not report sponsorship, but the corresponding author worked for Hill
	Dermaceuticals Inc, US.
	To be eligible for inclusion in the efficacy analysis, participants were permitted to deviate
	from the treatment plan $< = 2$ consecutive days and $< = 4/10$ days.
	No case of skin atrophy or telangiectasia were reported.
	This was a scalp trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	4.5%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Perez 1996

Methods	DESIGN
	Within-patient (1997)
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 84
T al ticipants	Treatment duration: 10 wks: FU: 52 wks
	LF: 0 (0%)
	BC: yes
	Age: 46 (range = 19 to 76)
	Gender (per cent men): 65.5%
	Severity: TSS (0 to 9) = 7.6 (range = 5 to 9)
	INCLUSION CRITERIA
	• Stable plaque or erythrodermic psoriasis

	 Unsatisfactory response to at least 1 previous treatment (topical steroids/UVB/ PUVA/MTX) Adult BSA ≥ 10% EXCLUSION CRITERIA Pregnant, nursing, or inadequate contraception Hepatic or renal impairment Recent systemic therapy or phototherapy or topical medications (excluding emollients)
Interventions	 Calcitriol 1.5 mcg/g OD (C) Placebo (vehicle) (P)
Outcomes	 Severity (erythema; plaque thickness; scaling) Total Severity Score (0 to 9) Investigator Global Assessment (5-pt, worse to excellent improvement) PASI (reported only for participants participating in follow-up study)
Notes	The NIH General Clinical Research Center supported the trial. There was also an uncontrolled follow up study (N = 22) involving large-area adminis- tration of calcitriol; 12-month results were based on results from 6 participants

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Pinheiro 1997

Methods	DESIGN
	Between-patient
	Participant delivery ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Open
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 132
I IIIII	Treatment duration: 8 wks; FU: 8 wks
	LF: 10 (7.6%)
	BC: yes
	Age: 48.2 (range = 17 to 90)
	Gender (per cent men): 59.1%
	Severity: per cent severe = 13.6%
	Duration (yrs): 16.9 (range = 0.5 to 60)
	INCLUSION CRITERIA
	Chronic plaque psoriasis
	• Adult
	• BSA $\geq 100 \text{ cm}^2$
	EXCLUSION CRITERIA
	Hypersensitivity to trial medicationsConcomitant treatment with Vitamin D/calcium/other relevant agent
	 Pregnancy
	Risk of pregnancy
	Lactation
	Unable to comply with protocol
Interventions	• Calcipotriol ointment 50 mcg/g BD (C)
	• Coal tar 5%/allantoin 2%/hydrocortisone cream 0.5% BD (T)
Outcomes	1. Signs (redness; thickness; scaliness)
	2. Total Sign Score (0 to 12)
	3. Investigator Global Assessment (5-pt: worse to cleared)
	4. Area of affected skin (area scales)
	5. Patient evaluation of overall response (VAS)
Notes	Leo Pharmaceuticals sponsored the trial.
	There was SD imputation (TSS).
Risk of bias	

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Pinheiro 1997 (Continued)

Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	7.6%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Poulin 2010

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT
	Described
Participants	 N: 217 Treatment duration: 6 mths; FU: 6 mths LF: 45 (20.7%) BC: yes Age: 50.4; range = 18 to 82 Gender (per cent men): 44.7% Ethnicity (per cent white): 91.2% Severity: GSS = 0 (clear): 10.1%; GSS = 0 (very mild): 33.7%; GSS = 0 (mild): 56.2%; < 20% scalp affected = 76.5%; mPASI = 4.2 (4.2SD) INCLUSION CRITERIA People aged > = 18 with moderate to severe scalp psoriasis (GSS = 3 or 4) eligible for initial phase (4 wks) Responders (GSS < 2) eligible for maintenance phase (6 mths) Topical and systemic wash-out periods required but not specified EXCLUSION CRITERIA Pregnancy or risk thereof Lactation Use of superpotent steroids on body Body psoriasis requiring systemic therapy.

Poulin 2010 (Continued)

Interventions	 Open-label initial treatment phase with clobetasol propionate 0.05% shampoo OD for up to 4 wks (N = 288), then participants with GSS < = 2 were randomised to maintenance therapy: clobetasol propionate 0.05% shampoo, 2 x/wk for 6 mths (CP) placebo (vehicle) shampoo, 2 x/wk for 6 mths (P) Participants evaluated every 4 wks for relapse (GSS > 2): relapses received clobetasol propionate 0.05% shampoo OD for 4 wks GSS < = 2: participants re-initiated maintenance therapy GSS > 2: participants exited the study
Outcomes	 Global Severity Score (GSS) (6-pt: 0 = clear to 5 = very severe) Extent of disease (6-pt, 0 (none) to 5 (80% to 100%)) (investigator) Signs (investigator): erythema, scaling, thickness (each scored 0 to 4) MPASI (0 to 7.2) Per cent with rebound at first relapse (mPASI > = 125% of baseline score) Pruritis (subject) (0 to 3) No. relapses Atrophy (0 to 2) Telangiectasia (0 to 2) Burning (0 to 2) adverse events HPA axis activity Patient satisfaction
Notes	Galderma Laboratories, LP, sponsored the trial. The trial (JDT) reported findings for a subset with moderate scalp psoriasis The sponsor supplied unpublished safety data. No useable efficacy data were available

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	20.7%
Baseline assessments	Low risk	These were reported (clinical and demo- graphic).
Baseline comparability demonstrated	Low risk	This was demonstrated.

Powers 2005

Risk of bias	Authors' indeement	Support for judgement	
Notes	Galderma Laboratories, LP, sponso The sponsor supplied unpublished This was an abstract only.		
Outcomes	success (clear/almost clear) and fai 2. Signs: erythema, scaling, thic 3. Dermatological Sum Score (I 4. Pruritus (0 to 4) 5. Investigator Global Improver 6. Subject Global Improvement 7. Routine clinical and safety lai	 Global Severity Score (GSS) (6-pt: 0 = none to 5 = very severe), dichotomised as success (clear/almost clear) and failure Signs: erythema, scaling, thickness (each scored 0 = absent to 4 = severe) Dermatological Sum Score (DSS) (0 to 12) Pruritus (0 to 4) Investigator Global Improvement score (7-pt: worse to clear) Subject Global Improvement score (7-pt: worse to clear) Routine clinical and safety laboratory parameters including calcium homeostasis (> 10% above upper limit of normal range) 	
Interventions	 Calcitriol 3 mcg/g ointment Placebo ointment BD (P) Max 30 g/day 	BD (C)	
Participants	N: 421 Treatment duration: 8 wks; FU: 8 LF: 0 (0%) BC: yes (clinically), demographics Age: NR Gender (per cent men): NR Per cent white: NR Severity: GSS (0 to 5) = 2.71 (0.4) INCLUSION CRITERIA • People aged > = 12 with mild • BSA < = 35% EXCLUSION CRITERIA • Not reported	NR (abstract) 5SD)	
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: unclear Concealment: unclear BLINDING Double-blind (participant/investig WITHDRAWAL/DROPOUT Not described		

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Powers 2005 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Unclear risk	Clinical assessments were reported (demo- graphic characteristics was unclear)
Baseline comparability demonstrated	Unclear risk	Clinical comparability was demonstrated (demographic comparability was unclear)

Reygagne 2005

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated randomisation list Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 151 Treatment duration: 4 wks; FU: 4 wks LF: 0 (0%) BC: yes Age: 45.3 (17.1SD)(range = 10 to 89) Gender (per cent men): 47.0% Per cent white: 98.7% Severity: TSS (0 to 9) = 4.90 (1.74SD); GSS (0 to 5) = 1.50 (0.60SD) Per cent scalp affected: 45% (SD28%) INCLUSION CRITERIA • People aged at least 12 with moderate to severe scalp psoriasis (GSS (0 to 5) > = 3 • Affected area on scalp > = 2 cm ² EXCLUSION CRITERIA • Very severe scalp psoriasis requiring systemic treatment • Known allergy to study medications • Immuno-compromised • History of adverse response to topical or systemic steroid therapy • Use of concomitant antipsoriatic therapy, betablockers, lithium, antimalarials or NSAIDs

Reygagne 2005 (Continued)

Interventions	 calcipotriol 0.005% solution BD (C) clobetasol propionate 0.05% shampoo OD (CP)
Outcomes	 Global Severity Score (GSS) (6-pt: 0 = none to 5 = very severe) Total Severity Score (10-pt: 0 = none to 9 = severe) Pruritus (0 to 3) Per cent scalp area affected Investigator's Global Assessment of Improvement (7-pt: worse to cleared) Subject's Global Assessment of Improvement (7-pt: worse to cleared) Atrophy, telangiectasia; burning sensation (scalp/neck/face) (0 to 3), burning/ stinging sensation (eyes) (0 to 3) Adverse events
Notes	Galderma Laboratories, LP, sponsored the trial. The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator).
Randomisation method reported	Low risk	A computer-generated randomisation list was used.
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Ruzicka 1998

OM plus Betamethasone valerate ointment ON 4 weeks (CB) Outcomes 1. PASI 2. Investigator Global Assessment (6-pt: deterioration to complete healing) 3. Patient evaluation of overall response (5-pt: scale NR) Notes The trial did not state sponsorship. Schering AG employed 1 author.	Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described
2. Calcipotriol 0.005% ointment BD 2 weeks, then calcipotriol ointment 0.005% OM plus Betamethasone valerate ointment ON 4 weeks (CB) Outcomes 1. PASI 2. Investigator Global Assessment (6-pt: deterioration to complete healing) 3. Patient evaluation of overall response (5-pt: scale NR) Notes The trial did not state sponsorship. Schering AG employed 1 author.	Participants	Treatment duration: 2 + 4 wks; FU: 14 wks LF: 7 (3.9%) BC: psoriasis comparable, demographics not reported Age: 42 (range = 18 to 80) Gender (per cent men): 55.6% Severity: PASI = 6.0 INCLUSION CRITERIA • Adults • Chronic plaque-type psoriasis • BSA \geq 30% • Calcium levels, renal, and liver function within normal range EXCLUSION CRITERIA • Pregnancy • Lactation
2. Investigator Global Assessment (6-pt: deterioration to complete healing) 3. Patient evaluation of overall response (5-pt: scale NR) Notes The trial did not state sponsorship. Schering AG employed 1 author.	Interventions	2. Calcipotriol 0.005% ointment BD 2 weeks, then calcipotriol ointment 0.005%
Schering AG employed 1 author.	Outcomes	2. Investigator Global Assessment (6-pt: deterioration to complete healing)
Risk of bias	Notes	
	Risk of bias	

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Ruzicka 1998 (Continued)

Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	3.9%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Salmhofer 2000

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: NR BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described
Participants	N: 63 Treatment duration: 4 wks; FU: 8 wks LF: 5 (7.9%) BC: yes Age: 47 (15.4SD, range = 19 to 83) Gender (per cent men): 54.0% Severity: PASI = 5.5 (2.65SD) Duration (months): 141 (124SD) INCLUSION CRITERIA • Stable chronic plaque psoriasis • Aged over 19 • Symmetrical lesions EXCLUSION CRITERIA • Other types of psoriasis • BSA affected > 30% • Concurrent systemic antipsoriatic therapy • Pregnancy • Lactation • Concurrent infectious disease • Other concurrent dematoses • Hypercalcaemia • Severe hepatic/renal disease
Interventions	 Calcipotriol ointment 5 mcg/g BD (C) Calcipotriol ointment 5 mcg/g OM plus diflucortolone valerate ointment 0.1%, ON (D)

Salmhofer 2000 (Continued)

Outcomes	 PASI IAGI (7-pt: extreme deterioration to complete healing) PAGI (7-pt: extreme deterioration to complete healing)
Notes	Schering Wien GmbH sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	7.9%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Sanchez 2001

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Unclear
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 28
	Treatment duration: 8 wks; FU: 8 wks
	LF: 3 (10.7%)
	LF: 3 (10.7%)
	LF: 3 (10.7%) BC: yes Age: 49.5 (range = 22 to 73)
	LF: 3 (10.7%) BC: yes Age: 49.5 (range = 22 to 73) Gender (per cent men): 50%
	LF: 3 (10.7%) BC: yes Age: 49.5 (range = 22 to 73)
	LF: 3 (10.7%) BC: yes Age: 49.5 (range = 22 to 73) Gender (per cent men): 50% Severity: PASI = 7.7 (range = 4 to 10)
	LF: 3 (10.7%) BC: yes Age: 49.5 (range = 22 to 73) Gender (per cent men): 50% Severity: PASI = 7.7 (range = 4 to 10) INCLUSION CRITERIA

Sanchez 2001 (Continued)

	 Systemic treatment within previous 4 wks Topical treatment within previous 2 wks Known hypersensitivity to sulphides Hypothyroidism Lactation
Interventions	Propylthiouracil cream 5% TDCalcipotriol ointment 50 mcg/g BD
Outcomes	1. PASI
Notes	The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	10.7%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Santoianni 2001

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated list using block randomisation; phar-
	macy-dispensed treatments in identical tubes
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 85
1 articipants	Treatment duration: 3 wks; FU: 3 wks
	LF: 4 (4.7%)

Santoianni 2001 (Continued)

	BC: clinical only Age: 51.9 (range = 18.8 to 88.5) Gender (per cent men): 55.3% Severity: PASI = 6.2 INCLUSION CRITERIA • Outpatients with disseminated keratotic plaque psoriasis • Aged over 18 EXCLUSION CRITERIA • Not reported
Interventions	Betamethasone 17-valerate 21-acetate plus tretinoine plus salicylic acid ODPlacebo OD
Outcomes	 PASI IAGI (5-pt: worse to cured) PAGI (5-pt: no change to excellent) Adverse events
Notes	IDI Farmaceuticia SpA sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	4.7%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Saraceno 2007

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-gen pensed sequentially Concealment: inadequate BLINDING Open WITHDRAWAL/DROPOUT Described	erated randomisation schedule; boxes dis-
Participants	 N: 150 Treatment duration: 12 wks; FU: 12 wks LF: 3 (2.0%) BC: yes Age: 47.7 (16.5SD); range = 18 to 83 Gender (per cent men): 66.0% Severity: per cent BSA = 18.6 (7.6SD), range 2 to 30; PASI = 9.2 (4.8SD), range = 1.6 to 33.0 Duration (yrs): 13.8 (13.4SD); range = 0 to 60.1 INCLUSION CRITERIA People aged > = 18 with mild to moderate plaque psoriasis EXCLUSION CRITERIA Topical therapy within previous 2 wks Systemic therapy within previous 4 wks Severe forms of plaque psoriasis Guttate, erythrodermic, and pustular psoriasis Cutaneous atrophy Suspected abnormality in calcium homeostasis Pregnancy Lactation 	
Interventions	 Calcipotriol 50 mcg/g cream BD (C) Calcipotriol 50 mcg/g plus betamethasone dipropionate 0.5 mg/g formulation (4 weeks) OD then calcipotriol 50 mcg/g cream (8 weeks) BD (C-B) 	
Outcomes	 PASI (scale NR) Skindex-29: burden of symptoms; social functioning; quality of life 	
Notes	Prodotti Formenti Srl, Milano, Italy, sponsored the trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	High risk	Consecutive coded treatment boxes were assigned to participants in a 1:1 ratio. It was

Saraceno 2007 (Continued)

		unclear if boxes were of identical appearance
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Low risk	A computer-generated randomisation schedule was used.
Loss to follow up	Low risk	2.0%
Baseline assessments	Low risk	These were reported (clinical and demo- graphic).
Baseline comparability demonstrated	Low risk	This was demonstrated (clinical and demo- graphic).

Scarpa 1994

Methods	DESIGN
	Between-patient
	Delivery unclear
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Blinding unclear
	WITHDRAWAL/DROPOUT
	Not described
D	
Participants	N: 160
	Treatment duration: 6 wks; FU: 10 wks
	LF: not reported
	BC: demographics comparable, severity not reported
	Age: 50
	Gender (per cent men): 68.1%
	Severity: not reported
	INCLUSION CRITERIA
	Plaque-type psoriasis
	EXCLUSION CRITERIA
	• Not reported
T	
Interventions	• Calcipotriol ointment 50 mg/g BD (C)
	• Betamethasone dipropionate ointment 0.05% + salicylic acid 3% BD (B)

Scarpa 1994 (Continued)

Outcomes	 Investigator Global Assessment (5-pt: null to excellent) Patient's overall acceptance (5-pt: null to excellent) 	
Notes	The trial did not report sponsorship.	

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Unclear risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Scarpa 1996

Methods	DESIGN Within-patient Delivery unclear ALLOCATION Random Method of randomisation: block randomisation (6 participants) Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 76 Treatment duration: 6 wks; FU: 8 wks LF: 13 (17.1%) BC: yes Age: not reported Gender (per cent men): not reported Severity: TSS (0 to 12) = 7.92 INCLUSION CRITERIA • Chronic plaque psoriasis EXCLUSION CRITERIA

Scarpa 1996 (Continued)

	 Concomitant medications (except emollients, tar shampoo, and salicylic acid) Topical or systemic steroids Calcium or vitamin D intake Antipsoriatic medications
Interventions	 Tacalcitol ointment 4 mcg/g OD (T) Betamethasone-17-valerate ointment 0.1% OD (B)
Outcomes	 Severity (erythema; thickness; scaling) Total Severity Score (0 to 12) Comparison of lesions, based on difference in TSS Investigator Global Assessment (6-pt: worsening to healing) Patient assessment of difference
Notes	The trial did not report sponsorship. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Block randomisation was used.
Loss to follow up	Low risk	17.1%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Scarpa 1997

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported; tubes labelled left or right and with participant ID number and tube ID number Concealment: unclear
	BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 157 Treatment duration: 6 wks; FU: 7 wks LF: 23 (14.6%) BC: yes Age: 49 (15SD; N = 134) Gender (per cent men): 65.6% (N = 157) Severity: TSS (0 to 12) = 7.7 INCLUSION CRITERIA • Stable chronic plaque psoriasis • Symmetrical lesions • In- and outpatients EXCLUSION CRITERIA • Pregnancy • Lactation • Inadequate contraception • Recent systemic, light, or topical therapy • Severe renal failure • Liver and cardiac dysfunction • Hypercalcemia • Hyperphosphoremia • AIDS • Drug addiction
Interventions	 Tacalcitol ointment 4 mcg/g OD (T) Placebo (vehicle) OD (P)
Outcomes	 Signs: scaling; erythema; scaling TSS (0 to 12) Patient compliance (tube count; tube contents)
Notes	The trial did not report sponsorship, but Istituto Gentili SpA provided medications and appeared to have undertaken the randomisation. There was SD imputation (TSS).

Risk of bias

Scarpa 1997 (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	14.6%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Sears 1997

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 190 participants Treatment duration: 3 wks; FU: 3 wks LF: 21 (11%) BC: yes Age: 44 (range = 19 to 73) Gender (per cent men): 47.9% Severity: TSS (0 to 9) = 6.0 Duration (yrs): 17 (range = 1 to 56) INCLUSION CRITERIA Mild or moderate psoriasis not spontaneously remitting Adults aged 18 to 70 Total Sign Score 3 to 8 EXCLUSION CRITERIA Acute systemic illness Hypothamic-pituitary-adrenal system disorder, severe hepatic or renal disorder Psoriatic infection Lactation, pregnancy, or inadequate contraception Recent use of any corticosteroid, long-acting antihistamines, retinoids

Sears 1997 (Continued)

	 Drugs exacerbating or influencing psoriasis Antimetabolic therapy PUVA ACE inhibitor Intolerant of topical corticosteroids or study medication
Interventions	 Hydrocortisone buteprate 0.1% cream BD (H) Placebo (vehicle) (P)
Outcomes	 Signs (erythema; skin thickening; scaling) Total Sign Score (0 to 9) Pruritis Investigator and patient evaluations of efficacy (4-pt: poor to excellent) Investigator Global Assessment (7-pt: exacerbation to cleared) Compliance (actual vs. expected usage)
Notes	The trial did not report sponsorship.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	11%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Seidenari 1997 (H)

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described
Participants	N: 14 Treatment duration: 6 wks; FU: 8 wks LF: 3 (21.4%) BC: yes Age: 46 (range 23 to 69, N = 26) Gender (per cent men): 46.2% (N = 26) Severity: TSS = 6.31 (1.25SD, N = 11) INCLUSION CRITERIA • Symmetrical, stable psoriatic plaques • Adult • Inpatient or outpatients EXCLUSION CRITERIA • Recent topical, UV, or systemic therapy • Inadequate contraception
Interventions	 Tacalcitol ointment 4 mcg/g OD (T) Betamethasone valerate ointment 0.1% OD (B)
Outcomes	 Signs: erythema; thickening; scaling Total Sign Score (0 to 12)
Notes	Demographic characteristics were summarised over both studies, placebo and active controls (N = 26)

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	21.4%

Seidenari 1997 (H) (Continued)

Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Seidenari 1997 (P)

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/assessor) WITHDRAWAL/DROPOUT Described	
Participants	N: 12 Treatment duration: 6 wks; FU: 8 wks LF: 1 (8%) BC: yes Age: 46 (range 23 to 69, N = 26) Gender (per cent men): 46.2% (N = 26) Severity: TSS = 6.50 (0.76SD, N = 11) INCLUSION CRITERIA • Symmetrical, stable psoriatic plaques • Adult • Inpatient or outpatients EXCLUSION CRITERIA • Recent topical, UV, or systemic therapy • Inadequate contraception	
Interventions	 Tacalcitol ointment 4 mcg/g OD (T) Placebo (vehicle) (P) 	
Outcomes	 Signs: erythema; thickening; scaling Total Sign Score (0 to 12) 	
Notes	Demographic characteristics were summarised over both studies, placebo and active controls (N = 26)	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.

Seidenari 1997 (P) (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/as- sessor).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	8.0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	-

Shuttleworth 1998

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: predetermined randomisation schedule in blocks of 10 Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 40 Treatment duration: 4 wks; FU: 4 wks LF: 3 (7.5%) BC: yes Age: 41.4 (12.0SD) Gender (per cent men):60.0% Severity: investigator's overall assessment (0 to 9) = 4.85; patient's overall assessment (0 to 4) = 2.43 INCLUSION CRITERIA Scalp psoriasis Aged 18 to 70 EXCLUSION CRITERIA Pregnancy Lactation Inadequate contraception Known hypersensitivity to study medication Participation in other study within previous month Concurrent systemic medication likely to affect psoriasis Photosensitivity; PUVA within previous 2 wks 'helmet' (diffuse) psoriasis Concurrent topical antipsoriatics Eye disease

Shuttleworth 1998 (Continued)

Interventions	Ciclopirox olamine shampoo 1.5% 3 times/wkPlacebo shampoo 3 times/wk
Outcomes	 IAGI (7-pt: very much worse to completely cleared) Area affected Investigator's overall assessment (extent of scalp psoriasis) (10-pt: normal to ≥ 75% scaling) Scaling (0 to 4.5) Pruritis Patient overall assessment (5-pt: poor to excellent) Adverse events
Notes	Stiefel Laboratories sponsored the trial. The study was underpowered to detect a statistically significant difference due to recruit- ment difficulties. This was a scalp trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Block randomisation was used.
Loss to follow up	Low risk	7.5%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Staberg 1989

Methods	DESIGN Within-patient Delivery unclear ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 10 Treatment duration: 6 wks; FU: 6 wks LF: 1 (10%) BC: yes Age: 50 (26 to 76) Gender (per cent men): not reported Severity: TSS (0 to 9) = 7.3 INCLUSION CRITERIA • Symmetrical chronic plaque psoriasis • Inpatients • Adult EXCLUSION CRITERIA • None reported	
Interventions	 Calcipotriol cream 1200 mcg/g BD (C) Placebo cream (P) 	
Outcomes	 Signs: infiltration; erythema; scaling Total Sign Score (0 to 9) 	
Notes	Leo Pharmaceutical Products supplied the drugs and undertook the statistical analysis	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	10.0%
Baseline assessments	Low risk	These were partially done.

Staberg 1989 (Continued)

Baseline comparability demonstrated	Low risk	-
Stein 2001		
Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: investigator undertook randomisation Concealment: inadequate BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 40 Treatment duration: 12 wks; FU: 12 wks LF: 3 (7.5%) BC: unclear Age: range = 20 to 70 + Gender (per cent men): not reported Severity: TSS (elbows) (0 to 12) = 7.0 INCLUSION CRITERIA • Mild to moderate symmetrical plaque psoriasis • Aged at least 18 EXCLUSION CRITERIA • Systemic treatment within previous 4 wks • Topical treatment within previous 2 wks • Investigational medication within previous 4 wks	
Interventions	 Betamethasone valerate foam 0.12% (Luxiq®) BD (B) Placebo foam BD (P) 	
Outcomes	 IAGI (7-pt: worse to completely clear) Composite severity score (sum of change scores in erythema, scaling, thickness) (0 to 12) Pruritis BSA involvement Compliance (weight of containers) 	
Notes	The Connetics Corporation sponsored the trial. There was SD imputation (IAGI).	
Risk of bias		
Bias	Authors' judgement	Support for judgement

Stein 2001 (Continued)

Allocation concealment (selection bias)	High risk	Allocation concealment was inadequate: The investigator undertook randomisation
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	7.5%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Stuecker 2001

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear
	BLINDING
	Blinding unclear
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 13 Treatment duration: 12 wks; FU: 12 wks LF: 2 (15.4%) BC: yes Age: 52.9 (12.2SD, range = 38 to 67) Gender (per cent men): 76.9% Severity: PASI = 9.11 (4.88SD, range = 2.20 to 18.70) Duration: 20.8 (12.7SD) INCLUSION CRITERIA • Stable psoriasis vulgaris • Aged 18 to 70 EXCLUSION CRITERIA • Topical treatment within previous wk • Modification of systemic treatment within previous 3 mths • Pototherapy within previous 6 wks • Known hypersensitivity to study medications • Avocado oil allergy • BSA $\geq 60\%$

Stuecker 2001 (Continued)

Interventions	 Calcipotriol cream BD Vitamin B cream (with avocado oil) BD
Outcomes	 PASI (modified to exclude head and neck; each side given weighting of 50%) IAGI (4-pt: poor to very good) PAGI (4-pt: poor to very good) Tolerability
Notes	Regeneratio Pharma AG sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	15.4%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Stutz 1996

Methods	DESIGN
	Within-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated randomisation code to allocate sides
	Concealment: adequate
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Not described
Participanto	N: 15
Participants	
	Treatment duration: 3 wks; FU: 3 wks
	LF: 2 (13.3%)
	BC: not reported

Stutz 1996 (Continued)

	Age: range = 21 to 68 Gender (per cent men): The trial did not report this. Severity: TSS (scale not reported) = 2.8 (0.3SD) INCLUSION CRITERIA • Mild plaque psoriasis EXCLUSION CRITERIA • Prescription treatments within previous 2 wks
Interventions	Polymyxin B cream 200,000 U/g TDPlacebo cream TD
Outcomes	 Total severity Erythema, scaling, thickness
Notes	Babcock Dermatologic Endowment and the alumni of the Department of Dermatology, University of Michigan Medical Center, MI, sponsored the trial

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	Body sites were selected then assigned a computer-generated randomisation code that was concealed from both investigators and participants
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	13.3%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Sudilovsky 1981

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: identical tubes allocated by random numbers table Concealment: adequate BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 78 (57% psoriasis) Treatment duration: 3 wks; FU: 3 wks LF: 0% BC: Inadequately reported Age: not reported Gender (per cent men): not reported Severity: not reported INCLUSION CRITERIA • Bilateral lesions of similar severity and duration EXCLUSION CRITERIA • Recent corticosteroid medication • History of poor response to corticosteroids • Concomitant local or systemic therapy that could affect psoriasis	
Interventions	 Halcinonide cream 0.1% OD + vehicle cream BD (H) Placebo (vehicle) treatment durations (P) 	
Outcomes	 Comparative therapeutic response (3-pt: equal response to markedly superior response) Absolute therapeutic response (4-pt: poor (< 25% improvement) to excellent (75% to 100% improvement)) Investigator Global Assessment: reflects comparative and absolute responses (methodology unclear) 	
Notes	The Squibb Institute for Medical Research sponsored the trial. Part of a larger study involving participants with atopic dermatitis and trialling other dosages; aggregated demographics were reported	
Risk of bias		
Bias	Authors' judgement	Support for judgement

Allocation concealment (selection bias) Low risk The part of the study (I or II) and side of the body chosen for treatment were concealed from investigators and determined by random numbers table	Dius	Hathors Judgement	Support for Judgement
	Allocation concealment (selection bias)	Low risk	the body chosen for treatment were con- cealed from investigators and determined

Sudilovsky 1981 (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Sutton 2001

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 53 participants Treatment duration: 8 wks; FU: 12 wks LF: 5 (9.4%) BC: yes Age: range = 26 to 67 Gender (per cent men): 58.5% Severity: duration (yrs) = 2 to 50 INCLUSION CRITERIA Psoriasis Good general health EXCLUSION CRITERIA Women of childbearing potential Systemic retinoids within previous 6 mths NSAIDs, cytostatic agents, or folic acid-containing vitamin preparations within previous mth Topical or UV treatments within previous 2 wks
Interventions	Methotrexate gel (Azone®) 1% ODPlacebo gel OD

Sutton 2001 (Continued)

Outcomes	 IAGI (6-pt: worse to cleared) Total Severity Score (erythema, thickness, scaling, pruritis) (0 to 20) Adverse events
Notes	Cato Research Ltd and Durham Pharmaceuticals LLC, NC, sponsored the trial. They treated lesions < = 20% BSA.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	9.4%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Syed 1996

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (participant/investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 60
i ur del purito	Treatment duration: 4 wks; FU: 52 wks
	LF: 0 (0%)
	BC: yes
	Age: 25.6 (range = 18 to 50)
	Gender (per cent men): 60.0%
	Severity: PASI = 9.3 (range = 4.8 to 16.7)
	Duration (yrs): 8.5 (range = 1 to 21)
	INCLUSION CRITERIA

Syed 1996 (Continued)

	 Mild-to-moderate chronic plaque-type psoriasis EXCLUSION CRITERIA Pregnancy Lactation Cytotoxic drugs Beta-blockers Recent systemic medication, UV therapy Epilepsy
Interventions	 Aloe vera extract 0.5% hydrophilic cream TDS (5 days/wk) Placebo cream TDS (5 days/wk)
Outcomes	 PASI Cure rate Significant clearing
Notes	The trial did not report sponsorship. Concomitant water soluble emollients were permitted. SDs were imputed (PASI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Syed 2001b

DESIGN		
Between-patient		
	orted	
Concealment: unclear		
BLINDING		
Double-blind (participant/investigation)	ator)	
Described		
N: 60 participants		
Treatment duration: 4 wks; FU: 8 v	wks	
-		
-	o 17.5)	
INCLUSION CRITERIA		
• Chronic, mild to moderate, plaque type psoriasis		
• Outpatients		
• PASI > 4 or BSA > 20%		
EXCLUSION CRITERIA		
• Topical or systemic corticosteroids or cytotoxic drugs or beta-blockers or		
· · · ·		
•		
 Account problems Concurrent renal, hepatic, or haematological abnormalities 		
,		
• Methotrexate gel (Azone ®) 0	0.25% BD (5 days/wk)	
• Placebo gel BD (5 days/wk)		
1. PASI		
2. Plaques cleared		
3. Adverse events		
4. Compliance (\leq 40 topical applications during 4-wk period)		
The trial did not report sponsorshi	p.	
SDs were imputation (PASI).	r.	
Authors' judgement	Support for judgement	
	Participant delivery ALLOCATION Random Method of randomisation: not rep Concealment: unclear BLINDING Double-blind (participant/investig WITHDRAWAL/DROPOUT Described N: 60 participants Treatment duration: 4 wks; FU: 8 LF: 0 (0%) BC: yes Age: 29.3 (range = 18 to 70) Gender (per cent men): 61.7% Severity: PASI = 9.8 (range = 5.3 tr Duration (yrs): 9.6 (range = 1 to 2 INCLUSION CRITERIA • Chronic, mild to moderate, p • Outpatients • PASI > 4 or BSA > 20% EXCLUSION CRITERIA • Topical or systemic corticoster phototherapy within previous 3 m • Pregnancy • Lactation • Alcoholic problems • Concurrent renal, hepatic, or • Methotrexate gel (Azone ®) (• Placebo gel BD (5 days/wk) 1. PASI 2. Plaques cleared 3. Adverse events	

Syed 2001b (Continued)

Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Tham 1994

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated random numbers Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 30 Treatment duration: 6 wks; FU: 6 wks LF: 3 (10%) BC: yes Age: 40 (range = 20 to 74) Gender (per cent men): 56.7% Ethnicity: Chinese (70.0%), Indian (16.7%), Malay (10.0%), and Sikh (3.3%) Severity: PASI = 6.65 Duration (years): 9.7 (range = 2 to 20) INCLUSION CRITERIA • Stable symmetrical chronic plaque-type psoriasis • Adult EXCLUSION CRITERIA • Recent systemic or UV therapy • Hypercalcaemia • High calcium or vitamin D intake • Impaired renal or hepatic function • Previous poor response to tar • Concomitant medications
Interventions	 Calcipotriol ointment 50 mcg/g BD (C) White soft paraffin OM plus coal tar solution BP in aqueous cream 15% ON (T)

Tham 1994 (Continued)

Outcomes	 PASI (modified to exclude head) Severity (erythema; infiltration; desquamation) Investigator Global Assessment (5-pt: worse to cleared) Patient Global Assessment (5-pt: worse to cleared)
Notes	Leo Pharmaceutical Products sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was single-blind (investigator).
Randomisation method reported	Low risk	Randomisation was computer-generated.
Loss to follow up	Low risk	10.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Tyring 2010

Methods	DESIGN Between-patient Participant delivery ALLOCATION
	Random Method of randomisation: computer-generated schedule (3:1), stratified by ethnicity;
	Concealment: adequate
	BLINDING
	Double-blind (participant/investigator) WITHDRAWAL/DROPOUT
	Described
Participants	N: 177 Treatment duration: 8 wks; FU: 52 wks LF: 0 (0%) (at 8 wks) BC: yes Age: 44.7 (range = 18 to 76) Gender (per cent men): 63.3% Ethnicity: per cent white = 0%; per cent Hispanic/Latino: 56%; per cent black/African American = 44% Severity: scalp IGA (moderate) = 80.2%; scalp IGA (severe/very severe) = 19.8%; TSS

	 (0 to 12) = 6.3, range = 4 to 11 Duration (yrs): 10.8, range = 1 to 50 INCLUSION CRITERIA People aged > = 18 with plaque psoriasis affecting the scalp and limbs/trunk > = 10% scalp affected IGA scalp psoriasis at least moderate Ethnicity: Hispanic/Latino/black/African American EXCLUSION CRITERIA Topical or UV therapy within previous 2 wks Biological within previous 12 wks
	 Systemic therapies within previous 4 wks Erythrodermic, exfoliative, or pustular psoriasis Skin infections Skin diseases confounding evaluation of psoriasis Calcium metabolic disorder or hypercalcaemia Pregnancy Lactation
Interventions	 Calcipotriene 50 mcg/g plus betamethasone 0.5 mg/g scalp formulation (gel) OD (C-B) Placebo gel OD (P) (8 wks) then C-B (44 wks)
	Maximum of 40 g/wk. Treatment was stopped if scalp psoriasis cleared Concurrent (scalp) antipsoriatics and 'chemical treatments of the hair' were not permitted All participants received C-B ointment for trunk/limbs.
Outcomes	 Investigator Global Assessment of Disease Severity (IGA): 6-pt = clear (0) to very severe (5). Based on written definitions for plaque thickness, scaling, and erythema. Success: IGA < = 1 Signs (investigator assessment): plaque thickness, scaling and erythema (each scored 0 (none) to 4 (very severe). Total Sign Score (0 to 12) Response criteria: TSS < = 1; PGA < = 1 Signs (each): absent Patient Global Assessment of Disease Severity (PGA): 6-pt = clear (0) to very severe (5). Based overall severity and impact on daily life Adverse events Blood pressure Laboratory tests: serum calcium, albumin, blood urea nitrogen, creatinine. Compliance (medication usage)
Notes	Leo Pharma A/S sponsored the trial This was part of a 52-wk study: Following 8 wks of double-blind treatment, all partici- pants received C-B scalp formulation for 44 wks. Data were reported only for the 8-wk end point The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	Participants were assigned the next consec- utive randomisation number available at the site for his/her ethnic category
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	Randomisation was preplanned according to a computer-generated randomisation schedule in a ratio of 3:1, stratified by eth- nicity
Loss to follow up	Unclear risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated (reported means, but not standard, deviations; pro- portions reported)

Tzung 2005

Methods	DESIGN
	Within-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not stated
	Concealment: unclear
	BLINDING
	Single-blind (investigator)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 23
Tarticipants	Treatment duration: 12 wks; FU: 16 wks
	LF: 4 (17.4%)
	BC: yes
	Age: 60.2 ; range = 12 to 80
	Gender (per cent men): 91.3%
	Severity: Signs = mean baseline scores ranged from 2.1 to 2.3 (SD range = 0.71 to 0.80)
	INCLUSION CRITERIA
	• People with plaque psoriasis attending study centre (veteran's hospital)
	Ethnicity: Chinese
	 Target lesions of at least moderate severity

	 EXCLUSION CRITERIA Topical within previous 2 wks Oral systemic therapies within previous 6 wks Phototherapy or sun exposure within previous 4 wks Unstable psoriasis Pregnancy Lactation Uncontrolled systemic disease
Interventions	 Calcipotriol 0.005% ointment BD (C) Tazarotene 0.1% gel ON placebo petrolatum ointment OM (T)
Outcomes	 Adverse events Scaling, plaque elevation, erythema (8 pt: 0 = none to 4 = very severe; 0.5 increments) Patient Assessment of Overall Improvement (PAGI): 5-pt = deterioration to excellent improvement (> 75%)
Notes	The trial did not state the sponsor. The number of adverse events was described, but the number of participants experiencing ADRs was not reported

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The study was single-blind (investigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	17.4%
Baseline assessments	Low risk	Clinical assessments were reported (within- patient study, so demographics compara- ble)
Baseline comparability demonstrated	Low risk	Clinical comparability was demonstrated.

Vali 2005

Vall 2009		
Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: flip of coin Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 42 Treatment duration: 8 wks; FU: 8 wks LF: 3 (7.1%) BC: yes Age: 32 (12.22SD) Gender (per cent men): 66.7% Severity: PASI = 10.86 (5.15SD) INCLUSION CRITERIA • People with symmetrical plaque psorie • BSA < 20% EXCLUSION CRITERIA • Systemic or topical antipsoriatic thera • Pregnancy • Lactation • Use of medications that could affect of	py within previous 4 wks
Interventions	 Topical caffeine 10% in Plastibase® T Placebo (Plastibase®) TD (P) 	ГD (C)
Outcomes	 modified PASI ('regional PASI'): rang Adverse events 	e unclear
Notes	The trial did not state the sponsor.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Randomisation was by the flip of a coin.
Loss to follow up	Low risk	7.1%

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Vali 2005 (Continued)

Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Van de Kerkhof 1989

Methods	DESIGNWithin-patientParticipant deliveryALLOCATIONRandomMethod of randomisation: not reportedConcealment: adequateBLINDINGDouble-blind (participant/investigator)WITHDRAWAL/DROPOUTDescribed
Participants	N: 10 Treatment duration: 4 wks; FU: 4 wks LF: 0 (0%) BC: yes Age: range = 28 to 72 Gender (per cent men): 70% Severity: TSS (0 to 9) = 7.2 INCLUSION CRITERIA • People with symmetrical chronic stable plaque psoriasis EXCLUSION CRITERIA • Topical antipsoriatic therapy within previous 2 wks • Systemic antipsoriatic therapy within previous 1 mth • BSA affected < = 15%
Interventions	 Calcitriol solution 2 mcg/ml BD (C) Placebo (vehicle) (P)
Outcomes	 Severity (erythema; thickness; scaling) TSS (0 to 9)
Notes	The trial did not report sponsorship, but Hoffmann-La Roche, Switzerland, employed 1 of the authors. The authors stated that allocation was concealed to the investigator, but provided no justification. Treatment was given at subtherapeutic dose. There was SD imputation (TSS).

Support for judgement Bias Authors' judgement

Van de Kerkhof 1989 (Continued)

Allocation concealment (selection bias)	Low risk	A randomisation code was generated cen- trally and concealed from the investigator until after trial completion
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Van de Kerkhof 1996a

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 122 Treatment duration: 8 wks; FU: 12 wks LF: 19 (15.6%) BC: inadequately reported Age: 44.8 (13.69SD) Gender (per cent men): 62.3% Severity: BSA = 5.6% Duration (mths): 233.5 (175.9SD) INCLUSION CRITERIA • Stable plaque psoriasis • Not localised on the scalp • BSA: 5.6% • Score ≥ 2 for erythema and desquamation and Score sum > 5 • White adults and adolescents EXCLUSION CRITERIA • Increased serum calcium or serum phosphate level • Recent systemic or topical antipsoriatic treatment • Serious disease

Van de Kerkhof 1996a (Continued)

	 Known allergy to study medication Recent participation in another clinical trial Expected poor compliance Calcium supplements Drugs influencing calcium metabolism Corticosteroids Barbiturates Phenytoin NSAIDs Pregnancy
Interventions	 Tacalcitol 4 mcg/g OD (T) Placebo (vehicle) (P)
Outcomes	 Signs: erythema; infiltration; desquamation Total Sign Score (0 to 12) Severity Preference assessment Area of test lesions Investigator Global Assessment (4-pt: poor to very good) Patient Global Assessment (4-pt: poor to very good) Assessment of benefit Post-treatment relapse
Notes	Hermal Kurt Herrmann sponsored the trial. Maximum treatment area: 10% BSA There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	15.6%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Van de Kerkhof 2002a

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Described
Participants	N: 88 participants Treatment duration: 4 wks; FU: 5 wks LF: 7 (8.0%) BC: yes Age: not reported Gender (per cent men): not reported Severity: PASI = 16.9 (range = 4.3 to 48.0) INCLUSION CRITERIA • Inpatients or outpatients • Aged 18 or over; chronic plaque psoriasis • Scalp involvement EXCLUSION CRITERIA • Other forms of psoriasis • Systemic antipsoriatic treatment within previous 6 wks • UV treatment within previous 6 wks • Impaired renal or hepatic function • History of urolithiasis or hypercalciurea • Arthritis • Immobilisation • Hypo- or hyperthyroidism • Heavy exposure to sunlight • Pregnancy • Lactation • Planning of pregnancy
Interventions	 Calcipotriol ointment 50 mcg/g (80 to 100 g/wk) and calcipotriol scalp solution 50 mg/ml (30 to 50 ml/wk) (C) Dithranol/tar regimen (D)
Outcomes	 PASI (scalp assessed separately) TSS (SCALP: erythema, thickness, scaling) (0 to 12) IAGI (6-pt: worse to clearance) PAGI (6-pt: worse to clearance) Adverse events
Notes	Leo Pharmaceutical Products sponsored the trial. The trial included inpatients. The trial reported medication quantities used: "patients complied well"

Van de Kerkhof 2002a (Continued)

'High dose' calcipotriol was given (above recommended dosage)
IAGI/PAGI: combined scalp/body
TSS: scalp only
PASI: body only

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	8.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Van de Kerkhof 2006

Methods	DESIGN Between-patient Nurse and participant delivery ALLOCATION Random Method of randomisation: computer-generated system using telephone response Concealment: adequate BLINDING Open WITHDRAWAL/DROPOUT Described
Participants	N: 106 Treatment duration: 12 wks; FU: 12 wks LF: 0 (0%) BC: yes Age (mean): 51.2; range = 25 to 83 Gender (per cent men): 59/100 (reported by de Korte 2008 - this is a secondary reference under Van de Kerkhof 2006); 6 participants missing) Ethnicity: NR Severity: mPASI (mean) = 9.9, range = 2.7 to 27.0 INCLUSION CRITERIA • Outpatients aged > = 18 with plaque psoriasis involving arms, trunk, legs, or a

	 combination of the aforementioned Condition amendable to topical treatment Target area treatable with < = 100 g calcipotriol/wk PASI extent score > = 2 (i.e. BSA (body region) > = 10%) Willing/able to comply with study protocol EXCLUSION CRITERIA Acute guttate, generalised pustular, or erythrodermic exfoliative psoriasis Atopic dermatitis, seborrhoeic dermatitis, or other inflammatory skin diseases Systemic antipsoriatic therapy (including corticosteroids) or phototherapy within previous 6 wks Topical antipsoriatic therapy (body) within previous 2 wks Planned changes in concurrent medication that could affect psoriasis (e.g. betablockers, lithium, etc) Pregnancy or risk thereof Lactation Known or suspected hypercalcaemia Hypersensitivity to components of study medication Treatment with an investigational drug within previous 3 mths Current participation in other clinical trial Planned exposure to excessive levels of sun/UV radiation Known treatment resistance to study medication
Interventions	 Calcipotriol 50 mcg/g ointment BD (C) Short-contact dithranol cream (dose titration from 0.1% to 5%; application time 15 to 45 mins) OD; class II or III corticosteroids for treatment of severe skin irritation (D) wk1: 5 (daily) visits to outpatient clinic wk2 onwards: twice-weekly visits PRN Participants intolerant of 0.1% dithranol used 0.05% dose. Cases of extensive hyperkeratosis, scales of participants in both treatment groups could be removed with salicylic acid (5%) in petrolatum before wk 0 Participants achieving clearance exited the study.
Outcomes	 PRIMARY OUTCOMES mPASI (scalp excluded) SECONDARY OUTCOMES IAGI: Investigator's overall assessment of treatment response: 6-pt (worse to clearance) PAGI: Patient's overall assessment of treatment response: 6-pt (worse to clearance) Quality of life Adverse events Compliance (per cent using study medication as prescribed)
Notes	Leo Pharma sponsored the trial. Secondary response criteria did not demonstrate a statistically significant difference be- tween treatments The sponsor supplied unpublished data.
Risk of bias	

Van de Kerkhof 2006 (Continued)

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	The treatment was assignment using a tele- phone voice response system, to ensure that the investigators' decision to randomise the participant preceded knowledge of the ran- domised treatment
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Low risk	Randomisation was through a computer- generated system using telephone response
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated (de Korte 2008). There were significantly more men in the dithranol group (72% versus 46%) . However, de Korte does not provide data on 6 participants. If all these participants are assumed to be of the gender that minimises between group differences, the difference is almost statistically significant at the 5% level (= 0.0502; Fisher's exact test). This may indicate a flawed randomisation process, or it may be chance. The primary reference does not report variance around mean PASI baseline estimates, so comparability was unclear

Van de Kerkhof 2009

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated schedule (1:2:2) Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 1417 Treatment duration: 8 wks; FU: 8 wks LF: 2 (0.1%) BC: yes Age: 48.3 (16.4SD); range = 18 to 92 Gender (per cent men): 44.8% Ethnicity (per cent white): 97.2% Severity: IGA (0 to 5) = 3.32 (0.71SD), range = 2 to 5; TSS (0 to 12) = 6.8 (1.8SD) Duration (yrs): 15.9 (13.4SD) INCLUSION CRITERIA • People aged > = 18 with scalp psoriasis involving > 10% of the scalp, amenable to treatment with < = 100 g medication/wk EXCLUSION CRITERIA • Concomitant topical scalp therapy (except medicated shampoos and emollients) • Planned initiation of/changes to concomitant medication that could affect scalp psoriasis • Use of topical treatment of the face, trunk or limbs with very potent (WHO group IV) corticosteroids or UVB within previous 2 wks • PUVA or grenz ray therapy, planned exposure to the sun, systemic therapy within previous 4 wks • Biological therapy within previous 6 mths • Erythrodermic, exfoliative, or pustular psoriasis • Presence of viral lesions, fungal, or bacterial skin infections, parasitic infections, or atrophic skin on the scalp, known/suspected abnormality of calcium homeostasis associated with clinically significant hypercalcaemia • Severe renal insufficiency • Severe hepatic disorders
Interventions	 Calcipotriol 50 mcg/g gel OD (C) Betamethasone dipropionate 0.5 mg/g gel OD (B) Calcipotriol 50 mcg/g plus betamethasone dipropionate 0. 5 mg/g gel (scalp formulation) OD (C-B) Participants achieving IGA = 0 could stop medication during the study
Outcomes	 Investigator's Global Assessment of severity of scalp psoriasis (IGA); 6-pt (0 = absence of disease to 5 = very severe disease) Responder: IGA absence or very mild Total Sign Score (erythema, thickness, scaliness): 13-pt (0 to 12)

Van de Kerkhof 2009 (Continued)

	 Patient assessment of overall response: 7-pt (worse to clear) Adverse events (local) Laboratory tests for calcium and albumin. Compliance (self report and weight medication used)
Notes	Leo Pharma A/S, Ballerup, Denmark, sponsored the trial. The sponsor supplied unpublished data.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	A computer-generated randomisation schedule was used (1:2:2)
Loss to follow up	Low risk	0.1%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Van der Vleuten 1995

Methods	DESIGN Within-patient Delivery unclear ALLOCATION Random Method of randomisation: not reported Concealment: unclear
	BLINDING Open WITHDRAWAL/DROPOUT Described
Participants	N: 10 Treatment duration: 2 wks; FU: 2 wks LF: 0 (0%) BC: inadequately reported Age: range = 20 to 72 Gender (per cent men): 40% Severity: PASI (modified) = 17.1 (2.1SEM)

Van der Vleuten 1995 (Continued)

	Duration (yrs): 3 to 53 INCLUSION CRITERIA • Adult • Inpatient • Severe, disabling psoriasis • Resistant to topical therapy EXCLUSION CRITERIA • Recent or concomitant oral antipsoriatic therapy, no topical or systemic treatments except corticosteroids for the scalp and face
Interventions	 Calcipotriol ointment 50 mcg/g BD Dithranol in paste or petroleum 0.05% to 4%, 24-hour application on alternate days
Outcomes	1. PASI (excludes scalp)
Notes	The trial did not report sponsorship. Treatment was delivered in an inpatient setting.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Vanderploeg 1976

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: identical tubes allocated by sequential admission number, corresponding to a standard randomisation schedule using a double-blind code Concealment: adequate BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 36 Treatment duration: 3 wks; FU: 3 wks LF: 3 of 36 (8.3%) BC: yes Age: 45.7 (range = 10 to 66; N = 33) Gender (per cent men): 48.5% (N = 33) Severity: TSS (0 to 20) = 9.9 INCLUSION CRITERIA • Psoriasis or atopic dermatitis EXCLUSION CRITERIA • Recent systemic or topical steroids • Concomitant medications
Interventions	 Betamethasone dipropionate ointment 0.05% BD (B) Vehicle BD (P)
Outcomes	 Signs: scale; erythema; pruritis; thickness; crusting Total Sign Score (0 to 20) Investigator Global Assessment (5 pt: exacerbation to excellent improvement)
Notes	The trial did not report sponsorship. This was part of a larger trial that included participants with atopic dermatitis (50% psoriasis). There was SD imputation (TSS).
Risk of bias	

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	The sequential trial admission number cor- responded to the randomisation schedule; the randomisation code was 'double blind'
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).

Vanderploeg 1976 (Continued)

Randomisation method reported	Low risk	Randomisation was sequential.
Loss to follow up	Low risk	8.3%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Veien 1997

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: block Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 287 Treatment duration: 8 wks; FU: 12 wks LF: 0 (0%) BC: psoriasis comparable, demographics inadequately reported Age: 45.0 Gender (per cent men): 53.7% Severity: BSA = 7.9% (range = 1% to 75%); TSS (0 to 12) = 7.59 INCLUSION CRITERIA • Adult • Stable plaque psoriasis • TSS > 5 • Erythema \geq 2, scaling \geq 2 EXCLUSION CRITERIA • Pregnancy • Lactation • High serum calcium, serum phosphate, serum creatinine • Unresponsive to calcipotriol • Intolerant to study ingredients • Serious co-morbidity
Interventions	 Tacalcitol ointment 4 mcg/g OD plus Tacalcitol vehicle OD (T) Calcipotriol ointment 50 mcg/g BD (C)
Outcomes	 Severity: erythema; infiltration; scaling; pruritus Total Sign Score (TSS): 0 to12 Investigator Global Assessment (6-pt: worse to clear)

Veien 1997 (Continued)

	 Patient Global Assessment (scale "virtually identical" to IAGI; details not reported) Patient evaluation of global usefulness (VAS) Patient evaluation of cosmetic acceptability Quantity of medication used Rebound (aggravation equal to or worse than pre-treatment severity)
Notes	Nycomed Pharma sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Block randomisation was used.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Vladimirov 1994

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT
Participants	Not described N: 60 Treatment duration: 6 wks; FU: 6 wks LF: 0 (0%) BC: inadequately reported Age: range = 18 to 70 Gender (per cent men): not reported Severity: PASI = 2.92

Vladimirov 1994 (Continued)

	Duration (yrs): range = 0.2 to 30 INCLUSION CRITERIA • Adult • Mild to moderate psoriasis EXCLUSION CRITERIA • None reported	
Interventions	 Calcipotriol cream 50 mcg/g BD (C) Betamethasone17-valerate ointment 0 	0.1% BD (B)
Outcomes	 PASI (range unclear) Investigator Global Assessment (scale NR) 	
Notes	Leo Pharmaceuticals sponsored the trial. There was SD imputation (PASI).	
Risk of bias		
Bias	Authors' judgement	Support for judgement

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Volden 1992

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described	
Participants	N: 10 Treatment duration: 4 wks; FU: 4 wks LF: 1 (10%) BC: yes Age: not reported Gender (per cent men): not reported Severity: BSA = 5% to 15% Duration (mean years): 20 INCLUSION CRITERIA • Symmetrical plaque-type psoriasis • Adult outpatients EXCLUSION CRITERIA • Recent active treatment for psoriasis	
Interventions	Dithranol 1% in petrolatum (D)Placebo (vehicle) (P)	
Outcomes	 Signs: erythema; infiltration; scaling Total Sign Score (0 to 12) 	
Notes	The trial did not report sponsorship, but Hydro Pharma, Swedenone, employed 1 of the authors	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.

Loss to follow up

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Low risk

10.0%

Volden 1992 (Continued)

Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-
Wall 1998		
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Not described	
Participants		
Interventions	Calcipotriol ointment 50Dithranol 0.1 to 2% OE	
Outcomes		

Wall 1998 (Continued)

Notes	Leo Pharmaceuticals sponsored the trial.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	7.2%
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Low risk	-

Weinstein 1996

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 324 Treatment duration: 12 wks; FU: 24 wks LF: 6 (1.9%) BC: yes Age: 46.8 (range = 12 to 83) Gender (per cent men): 67% Severity: per cent BSA = 6.9 (5.2SD); TSS (0 to 12) = 7.3 Duration (yrs): 17.5 (12.7SD) INCLUSION CRITERIA • Stable plaque psoriasis • BSA $\leq 20\%$ • 2 target lesions with plaque elevation ≥ 2 and ≥ 2 cm in diameter: 1 on elbow/ knee and 1 on trunk/limbs

Weinstein 1996 (Continued)

	EXCLUSION CRITERIA	
	Pustular or exfoliative psoriasis	
	Sensitivity to study medication	
	Other confounding skin conditions	
	• Recent use of tar shampoos	
	• Topical/systemic/light therapies	
	 Topical corticosteroids/UVB 	
	• PUVA/systemic therapy	
	Oral retinoids	
	Uncontrolled systemic disease	
	• Pregnant	
	• Lactating	
	Inadequate contraception	
Interventions	 Tazarotene gel 0.1% OD Tazarotene gel 0.05% OD Placebo (vehicle) (P) 	
Outcomes	1. Signs: plaque elevation; scaling; erythema	
Outcomes	2. Total Sign Score (0 to 12)	
	3. Per cent clearance	
	4. Patient assessment of cosmetic acceptability	
	1	
Notes	Allergan Inc. sponsored the trial.	
	The authors stated that concealment of treatment allocation was achieved for participants	
	and clinicians, as tubes were identical.	
	There was SD imputation (TSS).	

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	1.9%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Weinstein 2003

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: randomised in blocks of 6 Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT	
Participants	WITHDRAWAL/DROPOUTDescribedN: 1303Treatment duration: 12 wks; FU: 24 wksLF: 411 (31.5%)BC: yesAge: 48.2 (range = 18 to 84)Gender (per cent men): 62.6%Severity: OLA (0 to 5) (mean) = 3.6; BSA affected (mean) = 10.5%Duration (mean yrs): 18.4INCLUSION CRITERIA• Aged ≥ 18 • BSA $\geq 2\%$; OLA (0 to 5) ≥ 3 • Acceptable blood or urinary test resultsEXCLUSION CRITERIA• Pregnancy or risk thereof• Lactation• UV or topical therapies within previous 2 wks• PUVA or systemic therapies within previous 4 wks• Oral retinoid therapy within previous 8 wks• Expected prolonged exposure to UV light	
Interventions	 Tazarotene cream 0.05% OD (T1) Tazarotene cream 0.1% OD (T2) Placebo (P) 	
Outcomes	 Overall lesion assessment (OLA) (0 = none to 5 = very severe), as applied to all treated lesions Clinical success (OLA ≤ 2 at 12 wks) Effectiveness (improvement in OLA from baseline of ≥ 15% relative to placebo improvement score) Overall plaque elevation, scaling and erythema (each scored 0 = none to 4 = severe) Overall global response to treatment (7-pt: completely cleared to worsened) Target lesion response (7-pt: completely cleared to worsened) 	
Notes	Allergan Inc. sponsored the trial. The paper reported 2 trials; only study A reported follow-up data after 12 weeks (N = 108)	

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Low risk	Block randomisation was used.
Loss to follow up	High risk	31.5%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

White 2006 (H)

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: computer-generated schedule
	Concealment: unclear
	BLINDING
	Open (acute phase and maintenance phase for CB-W); double-blind (participant/inves-
	tigator) (maintenance phase for CB-C and CB-P)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 1136
1 u topuno	Treatment duration: 12 wks; FU: 12 wks
	LF: 0 (0%)
	BC: yes
	Age: 50.7; range = 18 to 89
	Gender (per cent men): 60.7%
	Ethnicity (per cent white): 96.9%
	Severity: per cent BSA = 12.1%; mPASI = 8.9 (3.8SD), range = 2.4 to 30.9; IGA moderate
	= 76%; IGA severe = 24%
	Duration (yrs): 0 to 75
	INCLUSION CRITERIA
	• People aged > = 18 with plaque psoriasis affecting at least 10% arms, 10% trunk,
	10% legs, or a combination of the aforementioned
	• IGA at least moderate
	EXCLUSION CRITERIA

White 2006 (H) (Continued)

	 Erythrodermic, exfoliative, pustular, or guttate psoriasis Skin infections Other confounding inflammatory skin disease Calcium metabolic disorder Pregnancy Lactation Concurrent antipsoriatic therapy
Interventions	 Calcipotriol 50 mcg/g plus betamethasone dipropionate 0.5 mg/g ointment OD (4 wks), then calcipotriol cream 50 mcg/g OD [8 wks] (CB-C) Calcipotriol 50 mcg/g plus betamethasone dipropionate 0.5 mg/g ointment OD (4 wks), then calcipotriol cream 50 mcg/g OD weekdays, CB ointment OD weekends (8 wks) (CB-W) Calcipotriol 50 mcg/g plus betamethasone dipropionate 0.5 mg/g ointment OD (4 wks), then placebo cream (calcipotriol vehicle), OD (8 wks) (CB-P)] Use < = 100 g/wk
Outcomes	 mPASI (0 to 64.8) - per cent change from baseline: rebound: > 125% of baseline; relapse: > 50% reduction in maximum improvement from baseline Investigator's Assessment of disease severity (IGA); 6-pt (absent to very severe) Patient's assessment of overall response (PAGI); 7-pt (worse to clear) Adverse events Compliance (medication usage; self-reported compliance)
Notes	Leo Pharma A/S sponsored the trial. Atrophy was not assessed.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open in the acute phase and maintenance phase for CB-W and dou- ble-blind (participant/investigator) in the maintenance phase for CB-C and CB-P. It was not possible to blind the alternating group (CB-W), as the vehicles were differ- ent
Randomisation method reported	Low risk	A preplanned computer-generated ran- domisation schedule was used (1:1:1)
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.

White 2006 (H) (Continued)

Baseline comparability demonstrated	Unclear risk	This was partially demonstrated (reported means, but not standard, deviations; pro- portions reported)
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White 2006 (P)

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated schedule Concealment: unclear BLINDING
	Open (acute phase and maintenance phase for CB-W); double-blind (participant/inves- tigator) (maintenance phase for CB-C and CB-P) WITHDRAWAL/DROPOUT Described
Participants	N: 1136 Treatment duration: 12 wks; FU: 12 wks LF: 0 (0%) BC: yes Age: 50.7; range = 18 to 89 Gender (per cent men): 60.7% Ethnicity (per cent white): 96.9% Severity: per cent BSA = 12.1%; mPASI = 8.9 (3.8SD), range = 2.4 to 30.9; IGA moderate = 76%; IGA severe = 24% Duration (yrs): 0 to 75 INCLUSION CRITERIA • People aged > = 18 with plaque psoriasis affecting at least 10% arms, 10% trunk, 10% legs, or a combination of the aforementioned • IGA at least moderate EXCLUSION CRITERIA • Erythrodermic, exfoliative, pustular, or guttate psoriasis • Skin infections • Other confounding inflammatory skin disease • Calcium metabolic disorder • Pregnancy • Lactation • Concurrent antipsoriatic therapy
Interventions	 Calcipotriol 50 mcg/g plus betamethasone dipropionate 0.5 mg/g ointment OD (4 wks), then calcipotriol cream 50 mcg/g OD [8 wks] (CB-C) Calcipotriol 50 mcg/g plus betamethasone dipropionate 0.5 mg/g ointment OD (4 wks), then calcipotriol cream 50 mcg/g OD weekdays, CB ointment OD weekends (8 wks) (CB-W) Calcipotriol 50 mcg/g plus betamethasone dipropionate 0.5 mg/g ointment OD

White 2006 (P) (Continued)

	(4 wks), then placebo cream (calcipotriol vehicle), OD (8 wks) (CB-P)] Use < = 100 g/wk
Outcomes	 mPASI (0 to 64.8) - per cent change from baseline: rebound: > 125% of baseline; relapse: > 50% reduction in maximum improvement from baseline Investigator's Assessment of disease severity (IGA); 6-pt (absent to very severe) Patient's assessment of overall response (PAGI); 7-pt (worse to clear) Adverse events Compliance (medication usage; self-reported compliance)
Notes	Leo Pharma A/S sponsored the trial. Atrophy was not assessed.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	High risk	The trial was open in the acute phase and maintenance phase for CB-W and dou- ble-blind (participant/investigator) in the maintenance phase for CB-C and CB-P. It was not possible to blind the alternating group (CB-W), as the vehicles were differ- ent
Randomisation method reported	Low risk	A preplanned computer-generated ran- domisation schedule was used (1:1:1)
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated (reported means, but not standard, deviations; pro- portions reported)

Wolska 1995

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described		
Participants	N: 52 Treatment duration: 4 wks; FU: 4 LF: 12 (23.1%) BC: yes Age: 44.7 (range = 18 to 77; N = Gender (per cent men): 70%; N Severity: TSS (0 to 9) (median) = INCLUSION CRITERIA • Plaque type psoriasis • At least 2 symmetrical lesion • TSS ≥ 6 • Between-plaque TSS scores • Lesions clinically stable ≥ 1 : EXCLUSION CRITERIA • Topical antipsoriatic treatment within previous 4 wks • UV treatment within previou • Pregnancy • Lactation; • Inadequate contraception	Treatment duration: 4 wks; FU: 4 wks LF: 12 (23.1%) BC: yes Age: 44.7 (range = 18 to 77; N = 40) Gender (per cent men): 70%; N = 40 Severity: TSS (0 to 9) (median) = 7.1 (range = 6.0 to 9.0) INCLUSION CRITERIA • Plaque type psoriasis • At least 2 symmetrical lesions, excluding those on neck, head, feet, and hands • TSS ≥ 6 • Between-plaque TSS scores ≤ 1 • Lesions clinically stable ≥ 1 wk EXCLUSION CRITERIA • Topical antipsoriatic treatment (tar/dithranol/steroids) within previous 2 wks • Systemic antipsoriatic treatment (steroids/retinoids/methotrexate/cyclosporin) within previous 4 wks • UV treatment within previous 4 wks • Pregnancy • Lactation;	
Interventions	Platelet aggregation activatinPlacebo solution BD	 Platelet aggregation activating factor (PAF) (Ro 24 to 0238) 10% solution BD Placebo solution BD 	
Outcomes	, , ,	 TSS (erythema, desquamation, infiltration) (0 to 9) IAGI (6-pt: marked worsening to total clearing) Adverse events 	
Notes		The trial did not report sponsorship. Inpatients PLS CAN YOU PUT THIS IN A SENTENCE?]	
Risk of bias			
Bias	Authors' judgement	Support for judgement	

Wolska 1995 (Continued)

Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/in- vestigator).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	23.1%
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Low risk	-

Wortzel 1975 (1)

Methods	DESIGN Between-patient Delivery unclear ALLOCATION Random Method of randomisation: sequential admission number Concealment: adequate BLINDING Double-blind (participant/physician) WITHDRAWAL/DROPOUT Not described
Participants	N: 76 Treatment duration: 3 wks; FU: 3 wks LF: 0 (0%) BC: not reported Age: not reported Gender (per cent men): not reported Severity: not reported INCLUSION CRITERIA • Moderately severe to very severe psoriasis and atopic dermatitis • Outpatients EXCLUSION CRITERIA • Not reported
Interventions	 Betamethasone dipropionate ointment 0.05 BD (B) Placebo BD (P)
Outcomes	 IAGI (5-pt: worse to excellent) Physician opinion of drug effect (scale unclear, results not reported)
Notes	Leo Pharmaceuticals sponsored the trial.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	The sequential trial admission number cor- responded to a treatment unit on a ran- domisation schedule
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/ physician).
Randomisation method reported	Low risk	Randomisation was sequential.
Loss to follow up	Low risk	0.0%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Wortzel 1975 (2)

Methods	DESIGN
inethous	Between-patient
	Delivery unclear
	ALLOCATION
	Random
	Method of randomisation: sequential admission number
	Concealment: adequate
	BLINDING
	Double-blind (participant/physician)
	WITHDRAWAL/DROPOUT
	Not described
Participants	N: 9
Tarticipanto	Treatment duration: 3 wks; FU: 3 wks
	LF: 0 (0%)
	BC: The trial did not report this.
	Age: not reported
	Gender (per cent men): not reported
	Severity: not reported
	INCLUSION CRITERIA
	• Moderately severe to very severe psoriasis and atopic dermatitis
	• Inpatients
	EXCLUSION CRITERIA
	Not reported

Wortzel 1975 (2) (Continued)

Interventions	 Betamethasone dipropionate ointment 0.05 BD (B) Placebo BD (P)
Outcomes	 IAGI (5-pt: worse to excellent) Physician opinion of drug effect (scale unclear, results not reported)
Notes	The trial did not report sponsorship. Concomitant water soluble emollients were permitted. There was SD imputation (PASI).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Low risk	The sequential trial admission number cor- responded to a treatment unit on a ran- domisation schedule
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (participant/ physician).
Randomisation method reported	Low risk	Randomisation was sequential.
Loss to follow up	Low risk	0.0%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Unclear risk	The trial did not report this.

Wozel 2001

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (unclear)
	WITHDRAWAL/DROPOUT
	Described
Participants	N: 38
I I I I I I I I I I I I I I I I I I I	Treatment duration: 2 wks; FU: 6 wks
	LF: 0 (0%)
	BC: yes (statistical significance not reported)

Wozel 2001 (Continued)

	Age: not reported Gender (per cent men): not reported Severity: not reported INCLUSION CRITERIA • Chronic plaque psoriasis EXCLUSION CRITERIA • Not reported	
Interventions	 Calcipotriol ointment OM plus fluocinolone acetonide ointment 0.025% ON (CF) Calcipotriol ointment OM plus vehicle ointment ON (CP) 4 weeks' maintenance for both groups with calcipotriol ointment BD 	
Outcomes	1. PASI	
Notes	The trial did not report sponsorship.	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient detail.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (it was unclear who was blinded).
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0.0%
Baseline assessments	Unclear risk	The trial did not report these.
Baseline comparability demonstrated	Low risk	-

Yang 2009

-	
Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Not stated WITHDRAWAL/DROPOUT Described
Participants	 N: 76 Treatment duration: 6 wks; FU: 6 wks LF: 0 (0%) BC: yes Age: range = 18 to 65 Gender (per cent men): 68.4% Severity: PASI = 14.3 (5.5SD), range = 4.2 to 19.6 Duration (yrs): range = 0.5 to 34 INCLUSION CRITERIA People aged 18 to 65 with plaque psoriasis affecting limbs and body PASI < 25 EXCLUSION CRITERIA Systemic antipsoriatic therapy within previous 4 wks Topical therapy within previous 2 wks Use of corticosteroids or vitamin D3 derivatives Concurrent infection Pregnancy Heart, liver, kidney, or mental disorder Known sensitivity to study medication External injury
Interventions	 Calcipotriol cream BD (C) Halometasone cream (OM), calcipotriol cream (ON) (2 wks); calcipotriol cream BD (weekdays), halometasone cream BD (weekends) (2 wks); calcipotriol cream BD (2 wks) (CH) Treatments were stopped when 90% improvement achieved.
Outcomes	 mPASI (0 to 64.8) Improvement score (based on change in PASI score x 100%) (IAGI): Complete recovery: > 90% Marked improvement: 60% to 89% Some improvement: 25% to 59% No effect: <25%
Notes	The trial did not state the sponsor. There was 1 case of skin atrophy in the monotherapy group. We received translation support for data extraction.

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Unclear risk	The trial did not report this.
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Low risk	0%
Baseline assessments	Low risk	These were reported.
Baseline comparability demonstrated	Low risk	This was demonstrated.

Zonneveld 1998 (H)

Methods	DESIGN Between-patient Delivery unclear ALLOCATION Random Method of randomisation: not reported Concealment: unclear BLINDING Double-blind (but unmatched regimens) (participant/assessor) WITHDRAWAL/DROPOUT Not described
Participants	N: 70 Treatment duration: 6 wks; FU: 6 wks LF: not reported BC: Yes (clinical); demographics not reported Age: not reported Gender (per cent men): not reported Severity: median LPSI ranged from 7.0 (C;T) to 8.0 (P) INCLUSION CRITERIA • Chronic plaque psoriasis • LPSI ≥ 6 EXCLUSION CRITERIA • Not reported
Interventions	 Calcipotriol ointment 0.005% BD (C) Tacrolimus ointment 0.3% OD (T) Placebo ointment BD (P)

Zonneveld 1998 (H) (Continued)

Outcomes	1. Local psoriasis severity index (scale NR)
Notes	The study was double-blind (but unmatched regimens). Fujisawa GmbH sponsored the trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (but unmatched regimens) (participant/assessor)
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	These were partially done.
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Zonneveld 1998 (P)

Methods	DESIGN
	Between-patient
	Delivery unclear
	ALLOCATION
	Random
	Method of randomisation: not reported
	Concealment: unclear
	BLINDING
	Double-blind (but unmatched regimens) (participant/assessor)
	WITHDRAWAL/DROPOUT
	Not described
Participants	N: 70
T articipants	Treatment duration: 6 wks; FU: 6 wks
	LF: not reported
	BC: yes (clinical), demographics not reported
	Age: not reported
	Gender (per cent men): not reported
	Severity: median LPSI ranged from 7.0 (C;T) to 8.0 (P)
	INCLUSION CRITERIA
	Chronic plaque psoriasis
	· · ·

Zonneveld 1998 (P) (Continued)

	 LPSI ≥ 6 EXCLUSION CRITERIA Not reported
Interventions	 Calcipotriol ointment 0.005% BD (C) Tacrolimus ointment 0.3% OD (T) Placebo ointment BD (P)
Outcomes	1. Local psoriasis severity index (scale NR)
Notes	The study was double-blind (but unmatched regimens). Fujisawa GmbH sponsored the trial. There was SD imputation (TSS).

Risk of bias

Bias	Authors' judgement	Support for judgement
Allocation concealment (selection bias)	Unclear risk	The trial reported insufficient details.
Blinding (performance bias and detection bias) All outcomes	Low risk	The trial was double-blind (but unmatched regimens) (participant/assessor)
Randomisation method reported	Unclear risk	The trial did not report this.
Loss to follow up	Unclear risk	The trial did not report this.
Baseline assessments	Low risk	-
Baseline comparability demonstrated	Unclear risk	This was partially demonstrated.

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Agrawal 2010	The study did not provide a comparison of interest (compared acitretin gel in 2 formulations)
Akerman 2009	The trialists did not report or make available adequate data
Ambroziak 2002	The study compared steroid against no treatment (rather than against placebo)
Baadsgaard 1995	The study was within-patient, but did not adopt clear left-right design and assessed multiple plaques for each participant

(Continued)

Baran 1999	The trialists did not report or make available adequate data
Bianchi 2003	The study compared steroid against no treatment (rather than against placebo)
Brodell 2011a	The sponsor did not report or make available adequate data.
Buder 2010	The study assessed multiple plaques for each participant (7 per participant)
Callen 1996	The trialists or sponsor did not report or make available adequate data
Cannavo 2003	This was a nail psoriasis trial, which was removed in the 2011 update
Carroll 2005	The comparator was not strictly placebo (salicylic acid in vehicle)
De Jong 1999	The comparator was not strictly placebo (urea in vehicle).
Elias 1994	It was unclear if it was a valid randomised controlled trial
Esposito 2009	The study did not provide a comparison of interest.
Friedrich 2004	The sponsors did not report or make available adequate data.
Hsia 2010	The study did not assess patient outcomes/tolerability
Insa 2009	The study did not assess patient outcomes/tolerability.
Iraji 2010	The trialists did not report or make available adequate data
Ito 2005	The study was not randomised.
Jansen 1986	No adequate data were reported.
Kaur 2004	The study was not randomised.
Kleyn 2005	The study did not provide a comparison of interest.
Kragballe 1989	Participants were randomised to the 2 substudies, but within the substudies, treatments were applied without randomisation
Kragballe 1994	This was a dose-ranging study of an unlicensed product not subsequently marketed
Lebwohl 1998b	The study did not provide a simple comparison against a vitamin D derivative treatment
Lebwohl 2001	The trialists or sponsor did not report or make available adequate data
Levin 2003	The trialists or sponsor did not report or make available adequate data

(Continued)

Meyrat 1996	The study did not provide a comparison of interest.
Palazon 2005	The study did not provide a comparison of interest.
Reygagne 2002a	The trialists or sponsor did not report or make available adequate data
Rhemus 2006	The study did not provide a comparison of interest (product unlicensed)
Ruzicka 2004	The trialists or sponsor did not report or make available adequate data
Sander 1998	The study did not provide a simple comparison against a vitamin D ³ derivative treatment
Saraswat 2007	The study did not provide a simple comparison against a vitamin D ⁴ derivative treatment
Scher 2001	This was a nail psoriasis trial, which was removed in the 2011 update
Sefton 1984	The trialists or sponsor did not report or make available adequate data
Sharma 2003	The study used concomitant UV light.
Syed 2001a	The trialists did not report or make available adequate data
Tokura 2004	No translation was available.
Tosti 1998	This was a nail psoriasis trial, which was removed in the 2011 update
Tzaneva 2003	The study used concomitant UV light.
van de Kerkhof 1996b	The trialists did not report or make available adequate data
Vena 2005	The study was not a randomised controlled trial.

Characteristics of studies awaiting assessment [ordered by study ID]

Bissonnette 2011

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: computer-generated code Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	 N: 61 Treatment duration: 12 wks; FU: 12 wks LF: 0 (0%) BC: yes Age: 50.6 (22 to 65) Gender (per cent men): 83.6% Severity: PGA = 3.3 (0.6SD); PASI = 6.0 (2.5SD); BSA = 3.2% (2.0SD) INCLUSION CRITERIA Participants aged 18 to 65 with mild to moderate plaque psoriasis BSA: 1% to 15% PGA: 2 to 4 Good general health EXCLUSION CRITERIA Concomitant serious illness/medical condition Systemic therapy within previous 12 to 24 wks (depending on drug) Phototherapy within previous 4 wks Topical antipsoriatic therapy within previous 2 wks Other non-plaque forms of psoriasis
Interventions	1% WBI-1001 in a cream formulation twice dailyPlacebo vehicle twice daily
Outcomes	 Physician's Global Assessment (PGA), 6-pt (0 = clear to 5 = very severe) Body surface area (BSA) Psoriasis Area and Severity Index (PASI)
Notes	Welichem Biotech Inc. sponsored the trial.

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Callis Duffin 2010

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Not described
Participants	 N: 200 Treatment duration: 12 wks; FU: 12 wks LF: NS BC: NS Age: NS Gender (per cent men): NS Severity: NS INCLUSION CRITERIA Participants aged 18 to 75 with mild to moderate plaque psoriasis BSA: 2% to 20% EXCLUSION CRITERIA Lesions solely involving intertriginious areas, the scalp, or the face Pustular psoriasis or erythroderma Concurrent systemic therapy, topical agents, or UVB therapy within 2 weeks of the first dose of study medication Systemic triazole antifungals except fluconazole
Interventions	INCB018424: Ruxolitinib phosphate cream QDPlacebo cream QD
Outcomes	 Total Lesion Score (sum of erythema, scaling, and thickness; points NS) PASI
Notes	Incyte Corporation sponsored the trial.

Calzavara-Pinton 2011

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Single-blind (investigator/outcome assessor) WITHDRAWAL/DROPOUT Described
Participants	N: 20 Treatment duration: 8 wks; FU: 16 wks LF: 2 (10%) BC: yes Age: 40.2 (range = 19 to 68) Gender (per cent men): 40% Severity: PSI = median 8 (IQR = 7 to 9.25) INCLUSION CRITERIA • Participants with mild plaque psoriasis • Symmetrical plaques EXCLUSION CRITERIA • Not stated
Interventions	 Budesonide 0.25 mg/g cream OM plus tacalcitol 4 mcg/g ointment ON Calcipotriol 50 mcg/g and betamethasone dipropionate 0.5 mg/g once daily
Outcomes	 Psoriasis Severity Index (erythema + infiltration + scaling) (0 to 12) VAS (adverse events; itching): 0 to 10 Patient preference and satisfaction (better/equal/worse)
Notes	The sponsor was not stated.

Henry 2011

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Single-blind (investigator) WITHDRAWAL Described
Participants	N: 13 Treatment duration: 4 wks; FU: 4 wks LF: 3 (23.0%) BC: NS Age: NS Gender (per cent men): NS Severity: NS INCLUSION CRITERIA • Participants with mild to moderate plaque psoriasis EXCLUSION CRITERIA • NS
Interventions	 Clobetasol propionate spray followed by calcitriol ointment twice daily Calcitriol ointment followed by clobetasol propionate spray twice daily
Outcomes	1. Signs: erythema, induration, scaling
Notes	Galderma sponsored the trial. This was an abstract only.

Katoh 2003

Methods	DESIGN
	Between-patient
	Participant delivery
	ALLOCATION
	Random
	Method of randomisation: not stated
	Concealment: unclear
	BLINDING
	Not stated
	WITHDRAWAL/DROPOUT
	Described
Danticinanta	N: 61
Participants	
	Treatment duration: 12 wks; FU: 12 wks
	LF: 2 (3%)

	BC: yes Age: 49.2 (14.3SD) Gender (per cent men): 34% Severity: PASI = 12.9 (5.3SD) Ethnicity: Japanese INCLUSION CRITERIA • Participants with stable plaque psoriasis EXCLUSION CRITERIA • Participants requiring systemic therapy or use of any antipsoriatic treatment within previous 4 wks
Interventions	 Calcipotriol 0.005% ointment once daily Combination treatment with calcipotriol 0.004% and 0.01% clobetasol propionate ointment once daily Treatments were applied after bathing. Participants achieving 50% reduction in baseline PASI then used calcipotriol once daily for a further 12 weeks
Outcomes	 Eruption score of trunk involvement: sum of erythema, scaling, and induration (0 to 12) PASI
Notes	The Japanese Ministry of Education, Science, Sports, and Culture sponsored the trial

Matheson 2011

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Open WITHDRAWAL/DROPOUT Not described
Participants	N: 32 Treatment duration: 2 wks; FU: 2 wks LF: NS BC: no Age: NS Gender (per cent men): NS Severity: NS INCLUSION CRITERIA • Participants aged \geq 12 with mild to moderate plaque psoriasis EXCLUSION CRITERIA • Not stated
Interventions	 Calcipotriol ointment 0.005% twice daily Calcipotriol foam 0.005% twice daily

Matheson 2011 (Continued)

Outcomes	1. Safety: adverse events; serum calcium
Notes	Stiefel, a GSK company, sponsored the trial.

Paulsen 2005

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 41 Treatment duration: 4 wks; FU: 12 wks LF: 1 (2%) BC: yes Age (median): 44 (23 to 77) Gender (per cent men): 65% Severity: duration (yrs; median) = 15 (range = 2 to 60) INCLUSION CRITERIA • Participants aged \geq 18 with stable mild to moderate plaque psoriasis • Test plaques similar within each participant (TSS \leq 1), TSS \leq 8 EXCLUSION CRITERIA • Concurrent use of topical steroids or vitamin D or emollients containing aloe vera • Pregnancy • Systemic or photo therapy within previous 2 mths • Topical antipsoriatic therapy within previous 2 wks • Known allergy to gel • Severe concomitant disease • Use of specific drugs
Interventions	Aloe vera gel twice dailyPlacebo gel twice daily
Outcomes	 PASI Local PASI (TSS) (0 to 9) Patient preference
Notes	Psoriasisfonden sponsored the trial.

Sadeghian 2011

Methods	DESIGN Between-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 60 Treatment duration: 12 wks; FU: 12 wks LF: 0 (0%) BC: yes Age: 32.5 (15.6SD) Gender (per cent men): 51.7% Severity: PASI = 3.9 (1.9SD) INCLUSION CRITERIA • Participants aged \geq 5 with plaque psoriasis • BSA \leq 10% EXCLUSION CRITERIA • Pregnancy • Non-plaque types of psoriasis • Participants with scalp and facial lesions
Interventions	Zinc pyrithione 0.25% cream twice dailyPlacebo cream twice daily
Outcomes	1. PASI (0 to 72)
Notes	The sponsor was not stated.

Vahlquist 2004

Methods	DESIGN Within-patient Participant delivery ALLOCATION Random Method of randomisation: not stated Concealment: unclear BLINDING Double-blind (participant/investigator) WITHDRAWAL/DROPOUT Described
Participants	N: 12 Treatment duration: 8 wks; FU: x wks LF: 0 (0%) BC: no Age: 43.8 (range = 21 to 58) Gender (per cent men): 83.3% Severity: NS INCLUSION CRITERIA • Participants with mild to moderate plaque psoriasis EXCLUSION CRITERIA • Systemic or topical therapy within previous 4 wks
Interventions	Thyroid hormone analogue (TriAc) 0.1% in hydrophilic ointment twice dailyPlacebo ointment twice daily
Outcomes	 Psoriasis Severity Index VAS to assess patient preference
Notes	-

Yang 1999

Methods	This will need to be translated; details to follow at next update
Participants	N: 60 Treatment duration: 6 wks; FU: 6 wks LF: unclear BC: unclear Age: range = 14 to 65 Gender (per cent men): unclear Severity: unclear INCLUSION CRITERIA • Unclear EXCLUSION CRITERIA • Unclear

Yang 1999 (Continued)

Interventions	Tacalcitol ointment17 alpha hydrocortisone cream
Outcomes	This will need to be translated; details to follow at next update
Notes	This will need to be translated; details to follow at next update

Characteristics of ongoing studies [ordered by study ID]

NCT00824980

Trial name or title	Combined Inhibition of Dipeptidyl Peptidase IV (DPIV/CD26) and Aminopeptidase N (APN/CD13) in the Treatment of Psoriasis - Phase II Single Center Study
Methods	Allocation: randomised End point classification: safety/efficacy study Intervention model: parallel assignment Masking: double-blind (participant, caregiver, investigator, outcomes assessor) Primary purpose: treatment
Participants	 Age 18 years of age at pre-study Diagnosis of plaque type psoriasis at least 3 month prior to enrolment Mild to moderate plaque type psoriasis with at least 2 plaques of approximately 15 cm² for which topical treatment is indicated
Interventions	Drug: IP10.C8 GelPlacebo Gel
Outcomes	1. Psoriasis Area and Severity Index at 4 weeks
Starting date	January 2009
Contact information	Alexander Narvarini, MD Telephone: +41 44 255 1111 alexander.navarini@usz.ch
Notes	Immune Technologies & Medicine GmbH http://clinicaltrials.gov/show/NCT00824980

Trial name or title	A Double-Blind, Vehicle-Controlled, Randomized, Dose Ranging, Multiple-Site Clinical Study to Evaluate
	the Efficacy and Safety of Desoximetasone Topical Sprays (0.05%, 0.25%) in Patients With Moderate to Severe Plaque Psoriasis
Methods	Allocation: randomised End point classification: safety/efficacy study Intervention model: parallel assignment Masking: double-blind (participant, caregiver, investigator, outcomes assessor) Primary purpose: treatment
Participants	 Have a definite clinical diagnosis of stable plaque psoriasis involving ≥ 10% of the body surface area (BSA) Have a combined Total Lesion Severity Score (TLSS) of ≥ 7 for the target lesion Have a plaque elevation score ≥ 3 of (moderate) for the target lesion The target lesion must have an area of at least 5 cm² Have a Physicians Global Assessment (PGA) score of 3 (moderate) or 4 (severe) at baseline for the overall disease severity
Interventions	 Drug: desoximetasone 0.05% once daily Drug: desoximetasone 0.05% twice daily Drug: desoximetasone 0.25% once daily Drug: desoximetasone 0.25% once daily Drug: vehicle once daily Drug: vehicle twice daily
Outcomes	 Clinical cure: Physician's Global Assessment (PGA) = 0 or 1 (time frame: 28 days) Lesion treatment success, score of 0 or 1, for each of the 3 signs of erythema, scaling, and plaque elevation (time frame: day 28)
Starting date	November 2009
Contact information	Darin B Brimhall, D.O., FACP Study Director Novum Pharmaceutical Research Services
Notes	Taro Pharmaceuticals USA http://clinicaltrials.gov/show/NCT01018134

NCT01206387

Trial name or title	A Double-Blind, Vehicle-Controlled, Randomized, Parallel Design, Multiple-Site Clinical Study to Evaluate the Efficacy and Safety of Desoximetasone 0.25% Topical Spray in Patients With Moderate to Severe Plaque Psoriasis
Methods	Allocation: randomised End point classification: safety/efficacy study Intervention model: parallel assignment Masking: double-blind (participant, investigator, outcomes assessor) Primary purpose: treatment

Participants	 Genders eligible for study: both Accepts healthy volunteers: no Men or non-pregnant, non-lactating women 18 years of age or older Have a definite clinical diagnosis of stable plaque psoriasis involving ≥10% of the body surface area (BSA) Have a combined Total Lesion Severity Score (TLSS) of ≥ 7 for the target lesion Have a plaque elevation score ≥ 3 of (moderate) for the target lesion The target lesion must have an area of at least 5 cm² Have a Physicians Global Assessment (PGA) score of 3 (moderate) or 4 (severe) at baseline for the overall disease severity
Interventions	Drug: desoximetasone spray 0.25%Drug: placebo comparator
Outcomes	 Clinical success (time frame: 28 days) - a participant is considered a clinical success if the Physician's Global Assessment (PGA) is 0 (clear) or 1 (almost clear) Treatment success (time frame: 28 days) - a participant is considered a treatment success for the target lesions if the target lesion has a score of 0 or 1 on the Target Lesion Severity Score (TLSS) for each of the 3 signs and symptoms (erythema, scaling, and plaque elevation) SECONDARY OUTCOMES Change in PGA (time frame: 28 days) - mean change from baseline to end of treatment in PGA
Starting date	August 2010
Contact information	Gail Gongas Telephone: 412 363 3300 ext: 522 GDGongas@novumprs.com
Notes	Taro Pharmaceuticals USA Study ID number: DSXS-0808 http://clinicaltrials.gov/show/NCT01206387
NCT01206660	

Trial name or title	A Double-Blind, Vehicle-Controlled, Randomized, Parallel Design, Multiple-Site Clinical Study to Evaluate the Efficacy and Safety of Desoximetasone 0.25% Topical Spray in Patients With Moderate to Severe Plaque Psoriasis
Methods	Study type: Interventional Study design: Allocation: randomised End point classification: safety/efficacy study Intervention model: parallel assignment Masking: double-blind (participant, investigator, outcomes assessor) Primary purpose: treatment

NCT01206660 (Continued)

Participants	Accepts healthy volunteers: no
	 Men or non-pregnant, non-lactating women 18 years of age or older
	• Have a definite clinical diagnosis of stable plaque psoriasis involving $\geq 10\%$ of the body surface area
	(BSA)
	• Have a combined Total Lesion Severity Score (TLSS) of ≥ 7 for the target lesion
	• Have a plaque elevation score \geq 3 of (moderate) for the target lesion
	• The target lesion must have an area of at least 5 cm ²
	• Have a Physicians Global Assessment (PGA) score of 3 (moderate) or 4 (severe) at baseline for the
	overall disease severity
Interventions	• Drug: desoximetasone spray 0.25%
interventions	 Drug: placebo
	• Drug. placebo
Outcomes	PRIMARY OUTCOMES
	1. Clinical success (time frame: 28 days) - a participant is considered a clinical success if the Physician's
	Global Assessment (PGA) is 0 (clear) or 1 (almost clear)
	2. Treatment success (time frame: 28 days) - a participant is considered a treatment success for the target
	lesions if the target lesion has a score of 0 or 1 on the Target Lesion Severity Score (TLSS) for each of the 3
	signs and symptoms (erythema, scaling, and plaque elevation)
	SECONDARY OUTCOMES
	1. Change in PGA (time frame: 28 days)
	2. Mean change from baseline to end of treatment in PGA
Starting date	August 2010
Contact information	Gail Gongas
Contact Information	Telephone: 412 363 3300 ext: 522
	GDGongas@novumprs.com
	GDGongas@novumprs.com
Notes	Taro Pharmaceuticals USA
	Study ID number: DSXS-0914
	http://clinicaltrials.gov/show/NCT01206660
NCT01246583	
Trial name or title	A Phase 2a, Multi Site, Randomized, Double Blind, Placebo Controlled, Parallel Group Study Of The Pilot
manne or trate	

	Efficacy, Safety, And Pharmacokinetics Of 2 Ointment Formulations Of CP-690,550 In Subjects With Mild To Moderate Chronic Plaque Psoriasis
Methods	Study type: interventional Study design: allocation: randomised End point classification: safety/efficacy study Intervention model: parallel assignment Masking: double-blind (participant, investigator) Primary purpose: treatment

Participants	Ages eligible for study: 18 years and older Genders eligible for study: both Accepts healthy volunteers: no INCLUSION CRITERIA • Mild to moderate chronic plaque psoriasis (psoriasis vulgaris), with the duration of at least 6 months • A target plaque of at least 9 sq. cm
	 EXCLUSION CRITERIA Demonstrates "rebound" or "flare" of chronic plaque psoriasis Non-plaque form of psoriasis Currently have or history of psoriatic arthritis Current drug-induced psoriasis Currently on systemic therapy or was on systemic therapy for psoriasis within the previous 6 months Currently on phototherapy for psoriasis or was on phototherapy within the previous 3 months
Interventions	 Drug: CP-690,550 ointment 1 Drug: vehicle 1 Drug: CP-690,550 ointment 2 Drug: vehicle 2
Outcomes	 PRIMARY OUTCOMES Per cent change from baseline at week 4 in Target Plaque Severity Score (TPSS) SECONDARY OUTCOMES Proportion of participants with Treatment Area Overall Severity of Psoriasis response of "clear" (0) or "almost clear" Proportion of participants with a difference from baseline of > = 2 steps in Treatment Area Overall Severity of Psoriasis score Per cent change from baseline at weeks 1, 2, 3, and 4 in target plaque area Change from baseline at weeks 1, 2, 3, and 4 in TPSS subscores for erythema, induration, and scaling Per cent change from baseline on the treatment area Itch Severity Item (ISI) Proportion of participants in each Patient Satisfaction with Study Medication (PSSM) response category Incidence, nature, and severity of observed and reported administration site adverse events over 4 weeks of treatment Incidence and severity of reactions of perilesional skin in the treatment area over 4 weeks of treatment Incidence of clinical laboratory abnormalities and change from baseline or 4 weeks of treatment Incidence of clinical laboratory abnormalities and change from baseline or 4 weeks of treatment Incidence of vital sign (blood pressure and heart rate) abnormalities and change from baseline in vital sign measures over 4 weeks of treatment Incidence of electrocardiogram (ECG) abnormalities and change from baseline in ECG measures over

NCT01246583 (Continued)

	4 weeks of treatmentPlasma CP-690,550 concentrations, from blood sampling
Starting date	November 22 2010
Contact information	Pfizer CT.gov Call Center
Notes	Pfizer

NCT01247818

Trial name or title	A Phase 2 Dose-Randomized, Vehicle-Controlled Study of PH-10-Aqueous Hydrogel for the Treatment of Plaque Psoriasis
Methods	Study type: interventional Study design: allocation: randomised End point classification: safety/efficacy study Intervention model: parallel assignment Masking: single-blind (outcomes assessor) Primary purpose: treatment
Participants	 Accepts healthy volunteers: no Men or women, age 18 or older Presence of mild to moderate plaque psoriasis on the trunk or extremities (excluding palms, soles, scalp, and facial or intertriginous areas) Fitzpatrick skin type I to VI Written informed consent by the participant or legal guardian
Interventions	 Drug: PH-10 (0.002% Rose Bengal) Drug: PH-10 (0.005% Rose Bengal) Drug: PH-10 (0.01% Rose Bengal) Drug: vehicle
Outcomes	 PRIMARY OUTCOMES The primary efficacy end point is "Treatment Success," a static end point assessed at day 29 after initial PH-10 treatment and defined as 0 or 1 on all Psoriasis Severity Index (PSI) components and 0 or 1 on the Plaque Response scale (time frame: 28 days) The primary safety end point is incidence of adverse experiences, including pain and dermatologic/ skin toxicity (incidence, severity, frequency, duration, and causality) (time frame: 8 weeks) SECONDARY OUTCOMES Psoriasis Severity Index (PSI) score changes at each visit from day 1 pre-treatment (time frame: 8 weeks) Plaque Response score changes at each visit from day 1 pre-treatment (time frame: 8 weeks) Pruritus Self-Assessment score changes at each visit from day 1 pre-treatment (time frame: 8 weeks)
Starting date	December 2010

NCT01247818 (Continued)

Contact information	Eric Wachter, PhD Provectus Pharmaceuticals
Notes	Provectus Pharmaceuticals http://clinicaltrials.gov/show/NCT01247818

NCT01422434						
Trial name or title	Efficacy and Safety of LEO 90105 Ointment (Calcipotriol Hydrate Plus Betamethasone Dipropionate) in Japanese Subjects With Psoriasis Vulgaris					
Methods	Study type: interventional Study design: allocation: randomised End point classification: efficacy study Intervention model: parallel assignment Masking: double-blind (participant, caregiver, investigator, outcomes assessor) Primary purpose: treatment					
Participants	 Accepts healthy volunteers: No Participants having understood and signed a written informed consent form prior to any study related procedures being carried out Japanese participants 20 years of age or above of either sex Clinical diagnosis of psoriasis vulgaris amenable to topical treatment, involving arms, trunk, legs, or any of the aforementioned A minimum mPASI score for extent of 2 in at least 1 body region (i.e. psoriasis affecting at least 10% of arms, 10% of trunk, 10% of legs, or any of the aforementioned) Psoriasis vulgaris on the trunk/limbs (excluding psoriasis on the genitals/skin folds) of not more than 30% body surface area (BSA) A target lesion of a minimum of 5 cm at its longest axis and preferably not located on an elbow or knee, scoring at least 3 for each of redness, thickness, and scaliness, and at least 10 in total by the physician's assessment of severity of the target lesion - a physician's global assessment of disease severity of psoriasis on trunk/limbs of mild, moderate, severe, or very severe Women of childbearing potential must have a negative result for a urine pregnancy test at day 0 (Visit 1) and must agree to use an adequate method of birth control, as judged by the (sub)investigator, during the study. The contraceptive method should have started an adequate amount of time before the pregnancy test, which is dependent on the particular method used and as judged by the (sub)investigator, and must continue for at least 1 week after the last application of study medication. A woman is defined as not of child-bearing potential if she is postmenopausal (12 months with no menses without an alter-native medical cause) or surgically sterile (tubal ligation/section, hysterectomy, or bilateral ovariectomy) 					
Interventions	 Drug: LEO 90105 = calcipotriol + betamethasone dipropionate Drug: Dovonex® = calcipotriol Drug: Rinderon® - DP = betamethasone dipropionate 					

NCT01422434 (Continued)

Outcomes	 PRIMARY OUTCOMES Change from baseline in Psoriasis Area and Severity Index (PASI) (time frame: week 0, week 4) SECONDARY OUTCOMES Change from baseline in target lesion assessment (time frame: week 0, week 4 Physician's global assessment of psoriasis (time frame: week 4) Adverse events (time frame: week 4)
Starting date	July 2011
Contact information	Akira Ozawa, MD, Professor, Principal Investigator Tokai University School of Medicine
Notes	LEO 90105 ointment contains both calcipotriol and betamethasone dipropionate. It has been approved for the treatment of psoriasis in more than 60 centres, including most European countries, the US, China, Korea, and Taiwan. This trial will investigate its safety and efficacy in the treatment of Japanese participants with psoriasis LEO Pharma http://clinicaltrials.gov/show/NCT01422434

NCT01465282

Trial name or title	A Randomized, Placebo-controlled, Phase IIb Study to Evaluate the Efficacy, Safety and Tolerability of 0. 05%, 0.1% and 0.5% w/w Topical CT327 When Applied Twice Daily in Subjects With Psoriasis Vulgaris					
Methods	Study type: interventional Study design: allocation: randomised End point classification: safety/efficacy study Intervention model: single group assignment Masking: double-blind (participant, caregiver, investigator, outcomes assessor) Primary purpose: treatment					
Participants	Accepts healthy volunteers: no INCLUSION CRITERIA • Men and women aged at least 18 years. • Stable psoriasis vulgaris EXCLUSION CRITERIA • Participants with guttate, erythrodermic, exfoliative, or pustular psoriasis.					
Interventions	 Drug: CT327 0.05% Drug: CT327 0.1% Drug: CT327 0.5% Drug: placebo 					
Outcomes	 PRIMARY OUTCOMES 1. Efficacy of CT327 ointment (0.05%, 0.1%, and 0.5% w/w) compared with placebo ointment SECONDARY OUTCOMES 1. Local and systemic toleration 					

NCT01465282 (Continued)

Starting date	November 1, 2011					
Contact information	No contacts or locations provided					
Notes	Creabilis SA					
NCT01536886						
Trial name or title	LEO 90100 Compared With Calcipotriol Plus Betamethasone Dipropionate Ointment, LEO 90100 Vehicle and Ointment Vehicle in Subjects With Psoriasis Vulgaris					
Methods	Allocation: randomised End point classification: efficacy study Intervention model: parallel assignment Masking: double-blind (caregiver, investigator, outcomes assessor) Primary purpose: treatment					
Participants	Accepts healthy volunteers: no					
	 INCLUSION CRITERIA Signed and dated informed consent obtained prior to any trial-related activities (including wash-out period) Age 18 years or above Either sex Any race or ethnicity All skin types Women of childbearing potential must have a negative pregnancy test at day 0 (Visit 1) Women of childbearing potential must agree to use a highly effective method of birth control during the study. A highly effective method of birth control is defined as 1 which results in a low failure rate (less than 1% per year) Able to communicate with the investigator and understand and comply with the requirements of the study. EXCLUSION CRITERIA Systemic treatment with biological therapies, whether marketed or not, with a possible effect on psoriasis vulgaris within the following time periods prior to randomisation: etanercept - within 4 weeks prior to randomisation other products - 4 weeks/5 half-lives (whichever is longer) Systemic treatment with all other therapies with a possible effect on psoriasis vulgaris (e.g. corticosteroids, retinoids, methotrexate, ciclosporin, and other immunosuppressants) within 4 weeks prior to randomisation Participants who have received treatment with any non-marketed drug substance (i.e. a drug which has not yet been made available for clinical use following registration) within 4 weeks/5 half-lives (whichever is longer) prior to randomisation PUVA therapy within 4 weeks prior to randomisation UVB therapy within 2 weeks prior to randomisation 					

NCT01536886 (Continued)

	 Planned excessive exposure of area(s) to be treated with study medication to either natural or artificial sunlight (including tanning booths, sun lamps, etc) during the study Planned initiation of, or changes to, concomitant medication that could affect psoriasis vulgaris (e.g. beta blockers, antimalarial drugs, lithium, ACE inhibitors) during the study Current diagnosis of guttate, erythrodermic, exfoliative, or pustular psoriasis Participants with any of the following conditions present on the treatment area: viral (e.g. herpes or varicella) lesions of the skin, fungal and bacterial skin infections, parasitic infections, skin manifestations in relation to syphilis or tuberculosis, acne vulgaris, atrophic skin, striae atrophicae, fragility of skin veins, icthyosis, ulcers, and wounds Other inflammatory skin disorders (e.g. seborrhoeic dermatitis or contact dermatitis) on the treatment area that may confound the evaluation of psoriasis vulgaris Known or suspected disorders of calcium metabolism associated with hypercalcaemia Known or suspected hypersensitivity to component(s) of the investigational products Current participation in any other interventional clinical study Previously randomised in this study Women who are pregnant, wishing to become pregnant during the study, or are breast-feeding
Interventions	 Drug: LEO 90100 Drug: betamethasone plus calcipotriol Drug: ointment vehicle Drug: LEO 90100 vehicle
Outcomes	1. Investigator's Global Assessment of Disease Severity (time frame: 4 weeks) Designated as safety issue: no
Starting date	May 2012
Contact information	Anders Ninn Hansen, MSc Pharm Telephone: +45 72 26 28 18 ext: 2818 anders.n-hansen@leo-pharma.com
Notes	-

NCT01536938

Trial name or title	LEO 90100 in the Treatment of Psoriasis Vulgaris
Methods	Study type: interventional Study design: allocation: randomised End point classification: efficacy study Intervention model: parallel assignment Masking: double-blind (participant, caregiver, investigator, outcomes assessor) Primary purpose: treatment
Participants	Accepts healthy volunteers: no INCLUSION CRITERIA • Signed and dated informed consent obtained prior to any trial related activities (including wash-out period)

- Age 18 years or above of either sex
- Any race or ethnicity
- All skin types
- Women of childbearing potential must have a negative pregnancy test at day 0 (Visit 1)

• Women of childbearing potential must agree to use a highly effective method of birth control during the study. A highly effective method of birth control is defined as one which results in a low failure rate (less than 1% per year)

• Able to communicate with the investigator and understand and comply with the requirements of the study

EXCLUSION CRITERIA

• Systemic treatment with biological therapies, whether marketed or not, with a possible effect on psoriasis vulgaris within the following time periods prior to randomisation:

- o etanercept within 4 weeks prior to randomisation
- o adalimumab, alefacept, infliximab within 8 weeks prior to randomisation
- o ustekinumab within 16 weeks prior to randomisation
- o other products 4 weeks/5 half-lives (whichever is longer)
- Systemic treatment with all other therapies with a possible effect on psoriasis vulgaris (e.g. corticosteroids, retinoids, methotrexate, ciclosporin, and other immunosuppressants) within 4 weeks prior

to randomisation

• Participants who have received treatment with any non-marketed drug substance (i.e. a drug which has not yet been made available for clinical use following registration) within 4 weeks/5 half-lives (whichever is longer) prior to randomisation

- PUVA therapy within 4 weeks prior to randomisation
- UVB therapy within 2 weeks prior to randomisation

• Planned excessive exposure of area(s) to be treated with study medication to either natural or artificial sunlight (including tanning booths, sun lamps, etc) during the study

• Planned initiation of, or changes to, concomitant medication that could affect psoriasis vulgaris (e.g. beta blockers, antimalarial drugs, lithium, ACE inhibitors) during the study

• Current diagnosis of guttate, erythrodermic, exfoliative, or pustular psoriasis

• Participants with any of the following conditions present on the treatment area: viral (e.g. herpes or varicella) lesions of the skin, fungal and bacterial skin infections, parasitic infections, skin manifestations in relation to syphilis or tuberculosis, acne vulgaris, atrophic skin, striae atrophicae, fragility of skin veins, icthyosis, ulcers, and wounds

• Other inflammatory skin disorders (e.g. seborrhoeic dermatitis or contact dermatitis) on the treatment area that may confound the evaluation of psoriasis vulgaris

- Known or suspected disorders of calcium metabolism associated with hypercalcaemia
- Known or suspected severe renal insufficiency or severe hepatic disorders
- Known or suspected hypersensitivity to component(s) of the investigational products
- Current participation in any other interventional clinical study
- Previously randomised in this study

Interventions	 Drug: LEO 90100 Drug: calcipotriol Drug: betamethasone
Outcomes	1. Investigator's Global Assessment of Disease Severity (time frame: 4 weeks) (Designated as safety issue: no)
Starting date	May 2012

Topical treatments for chronic plaque psoriasis (Review)

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NCT01536938 (Continued)

Contact information	Anders Ninn Hansen, MSc Pharm Telephone: +45 72 26 28 18 ext: 2818 anders.n-hansen@leo-pharma.com
Notes	-

DATA AND ANALYSES

Comparison 1. Vitamin D analogues versus placebo

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	20		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol	10	2287	Std. Mean Difference (IV, Random, 95% CI)	-0.93 [-1.17, -0.68]
1.2 Calcipotriol plus occlusion	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.3 Calcitriol	6	1120	Std. Mean Difference (IV, Random, 95% CI)	-1.03 [-1.71, -0.36]
1.4 Tacalcitol	2	433	Std. Mean Difference (IV, Random, 95% CI)	-0.84 [-1.41, -0.26]
1.5 Maxacalcitol	1	103	Std. Mean Difference (IV, Random, 95% CI)	-1.43 [-1.91, -0.96]
1.6 Paricalcitol OD	1	22	Std. Mean Difference (IV, Random, 95% CI)	-1.66 [-2.66, -0.67]
1.7 Becocalcidiol OD	1	121	Std. Mean Difference (IV, Random, 95% CI)	-0.22 [-0.58, 0.14]
1.8 Becocalcidiol twice daily	1	119	Std. Mean Difference (IV, Random, 95% CI)	-0.67 [-1.04, -0.30]
2 TSS	19		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 Calcipotriol	10	1208	Std. Mean Difference (IV, Random, 95% CI)	-1.15 [-1.41, -0.89]
2.2 Calcipotriol plus occlusion	1	187	Std. Mean Difference (IV, Random, 95% CI)	-0.15 [-0.44, 0.14]
2.3 Calcitriol	4	1027	Std. Mean Difference (IV, Random, 95% CI)	-1.22 [-2.38, -0.07]
2.4 Tacalcitol	3	496	Std. Mean Difference (IV, Random, 95% CI)	-0.66 [-0.95, -0.36]
2.5 Maxacalcitol	1	103	Std. Mean Difference (IV, Random, 95% CI)	-1.61 [-2.10, -1.12]
2.6 Paricalcitol OD	1	22	Std. Mean Difference (IV, Random, 95% CI)	-2.15 [-3.24, -1.06]
2.7 Becocalcidiol OD	1	121	Std. Mean Difference (IV, Random, 95% CI)	-0.02 [-0.37, 0.34]
2.8 Becocalcidiol twice daily	1	119	Std. Mean Difference (IV, Random, 95% CI)	-0.46 [-0.83, -0.10]
3 PASI	9	2422	Std. Mean Difference (IV, Random, 95% CI)	-0.58 [-0.71, -0.45]
3.1 Calcipotriol	8	2195	Std. Mean Difference (IV, Random, 95% CI)	-0.65 [-0.75, -0.55]
3.2 Calcipotriol plus occlusion	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
3.3 Calcitriol	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
3.4 Tacalcitol	1	227	Std. Mean Difference (IV, Random, 95% CI)	-0.27 [-0.56, 0.03]
3.5 Maxacalcitol	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
3.6 Paricalcitol OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.7 Becocalcidiol OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.8 Becocalcidiol twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	5	1467	Std. Mean Difference (IV, Random, 95% CI)	-0.54 [-0.72, -0.36]
4.1 Calcipotriol	2	439	Std. Mean Difference (IV, Random, 95% CI)	-0.64 [-0.97, -0.30]
4.2 Calcipotriol plus occlusion	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Calcitriol	2	801	Std. Mean Difference (IV, Random, 95% CI)	-0.59 [-0.76, -0.41]
4.4 Tacalcitol	1	227	Std. Mean Difference (IV, Random, 95% CI)	-0.24 [-0.53, 0.05]
4.5 Maxacalcitol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.6 Paricalcitol OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.7 Becocalcidiol OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.8 Becocalcidiol twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	30		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 Calcipotriol	17	3269	Std. Mean Difference (IV, Random, 95% CI)	-0.96 [-1.15, -0.77]
5.2 Calcipotriol plus occlusion	1	187	Std. Mean Difference (IV, Random, 95% CI)	-0.15 [-0.44, 0.14]
5.3 Calcitriol	7	1140	Std. Mean Difference (IV, Random, 95% CI)	-0.92 [-1.54, -0.29]
5.4 Tacalcitol	4	723	Std. Mean Difference (IV, Random, 95% CI)	-0.73 [-1.09, -0.37]
5.5 Maxacalcitol	1	103	Std. Mean Difference (IV, Random, 95% CI)	-1.43 [-1.91, -0.96]

5.6 Paricalcitol OD	1	22	Std. Mean Difference (IV, Random, 95% CI)	-1.66 [-2.66, -0.67]
5.7 Becocalcidiol OD	1	121	Std. Mean Difference (IV, Random, 95% CI)	-0.22 [-0.58, 0.14]
5.8 Becocalcidiol twice daily	1	119	Std. Mean Difference (IV, Random, 95% CI)	-0.67 [-1.04, -0.30]
6 Total withdrawals	25	4715	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.05, 0.00]
6.1 Calcipotriol	14	3132	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.06, 0.00]
6.2 Calcipotriol plus occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.3 Calcitriol	5	339	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.02]
6.4 Tacalcitol	4	857	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.14, 0.05]
6.5 Maxacalcitol	1	120	Risk Difference (M-H, Random, 95% CI)	0.11 [-0.01, 0.23]
6.6 Paricalcitol OD	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
6.7 Becocalcidiol OD	1	124	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.12, 0.11]
6.8 Becocalcidiol twice daily	1	121	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.13, 0.10]
7 Withdrawals due to adverse	23	4463	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.01]
events	23	4403	Risk Difference (NI-11, Randolli, 9970 CI)	-0.00 [-0.02, 0.01]
7.1 Calcipotriol	12	2880	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.04, 0.00]
7.2 Calcipotriol plus occlusion	0	2000	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.3 Calcitriol	5	339	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.02]
7.4 Tacalcitol	4	857	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.01, 0.02]
7.5 Maxacalcitol	1	120	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.08, 0.06]
7.6 Paricalcitol OD	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
7.7 Becocalcidiol OD	1	124	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.02, 0.08]
7.8 Becocalcidiol twice daily	1	124	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.01, 0.11]
8 Withdrawals due to treatment	14	2752	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.05, 0.00]
failure	14	27 32	Kisk Difference (M-11, Kandolit, 99% CI)	-0.05 [-0.05, 0.00]
8.1 Calcipotriol	7	1770	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.08, 0.01]
8.2 Calcipotriol plus occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.3 Calcitriol	4	281	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.12, 0.05]
8.4 Tacalcitol	1	314	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
8.5 Maxacalcitol	1	120	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.11, 0.04]
8.6 Paricalcitol OD	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
8.7 Becocalcidiol OD	1	124	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.07, 0.04]
8.8 Becocalcidiol twice daily	1	121	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.09, 0.02]
9 Adverse events (local)	19	4402	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.01, 0.02]
9.1 Calcipotriol	11	2652	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.03, 0.05]
9.2 Calcipotriol plus occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.3 Calcitriol	4	917	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.03, 0.02]
9.4 Tacalcitol	2	566	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.03, 0.03]
9.5 Maxacalcitol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.6 Paricalcitol OD	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
9.7 Becocalcidiol OD	1	124	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.06, 0.08]
9.8 Becocalcidiol twice daily	1	121	Risk Difference (M-H, Random, 95% CI)	0.10 [0.00, 0.19]
10 Adverse events (systemic)	14	2463	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.1 Calcipotriol	8	1182	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.2 Calcipotriol plus	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
occlusion	0	Ũ		010 [010, 010]
10.3 Calcitriol	3	647	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.4 Tacalcitol	1	244	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
10.5 Maxacalcitol	1	120	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
10.6 Paricalcitol OD	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
10.7 Becocalcidiol OD	1	124	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
10.8 Becocalcidiol twice daily	1	124	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
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Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	11	1904	Std. Mean Difference (IV, Random, 95% CI)	1.00 [-1.18, -0.82]
1.1 Betamethasone	2	739	Std. Mean Difference (IV, Random, 95% CI)	-0.81 [-0.98, -0.64]
dipropionate OD				
1.2 Betamethasone	4	537	Std. Mean Difference (IV, Random, 95% CI)	-1.35 [-1.56, -1.15]
dipropionate twice daily				
1.3 Betamethasone	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
dipropionate, maintenance				
1.4 Betamethasone valerate	1	74	Std. Mean Difference (IV, Random, 95% CI)	-1.41 [-1.92, -0.90]
1.5 Budesonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.6 Desonide	1	76	Std. Mean Difference (IV, Random, 95% CI)	-0.81 [-1.34, -0.28]
1.7 Diflorasone diacetate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.8 Fluticasone propionate	2	383	Std. Mean Difference (IV, Random, 95% CI)	-0.93 [-1.14, -0.72]
1.9 Hydrocortisone buteprate	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
1.10 Mometasone furoate	1	95	Std. Mean Difference (IV, Random, 95% CI)	-0.75 [-1.17, -0.34]
2 TSS	7	611	Std. Mean Difference (IV, Random, 95% CI)	-0.77 [-1.01, -0.52]
2.1 Betamethasone dipropionate OD	1	93	Std. Mean Difference (IV, Random, 95% CI)	-0.74 [-1.16, -0.32]
2.2 Betamethasone	1	33	Std. Mean Difference (IV, Random, 95% CI)	-0.77 [-1.48, -0.06]
dipropionate twice daily	1	55	Std. Wear Difference (17, Randolli, 7976 Ci)	0.77 [1.10, 0.00]
2.3 Betamethasone	1	38	Std. Mean Difference (IV, Random, 95% CI)	-0.95 [-1.62, -0.27]
dipropionate, maintenance	1	50	Std. Wear Difference (1V, Randolli, 7570 Cl)	-0.77 [-1.02, -0.27]
2.4 Betamethasone valerate	1	22	Std. Mean Difference (IV, Random, 95% CI)	-1.09 [-2.00, -0.18]
2.5 Budesonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.6 Desonide	1	76	Std. Mean Difference (IV, Random, 95% CI)	-1.16 [-1.70, -0.61]
2.7 Diflorasone diacetate	1	93	Std. Mean Difference (IV, Random, 95% CI)	-0.32 [-0.73, 0.09]
2.8 Fluticasone propionate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.9 Hydrocortisone buteprate	1	161	Std. Mean Difference (IV, Random, 95% CI)	-0.46 [-0.77, -0.15]
2.10 Mometasone furoate	1	95	Std. Mean Difference (IV, Random, 95% CI)	-1.12 [-1.55, -0.68]
3 PASI	3	1158	Std. Mean Difference (IV, Random, 95% CI)	-0.97 [-1.31, -0.62]
3.1 Betamethasone dipropionate OD	2	739	Std. Mean Difference (IV, Random, 95% CI)	-0.79 [-1.44, -0.14]
3.2 Betamethasone	1	419	Std. Mean Difference (IV, Random, 95% CI)	-1.21 [-1.44, -0.97]
dipropionate twice daily	-	>		
3.3 Betamethasone	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
dipropionate, maintenance	Ũ	0		010 [010, 010]
3.4 Betamethasone valerate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.5 Budesonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.6 Desonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.7 Diflorasone diacetate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.8 Fluticasone propionate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.9 Hydrocortisone buteprate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.10 Mometasone furoate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.1 Betamethasone dipropionate OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

4.2 Potemathecone	0	0	Std Moon Difference (IV Dendem 050/ CI)	0 0 [0 0 0 0]
4.2 Betamethasone dipropionate twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Betamethasone	0	0	Std Mars Difference (IV Denderer 050/ CI)	0.0.[0.0.0.0]
4.5 Betamethasone dipropionate, maintenance	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.4 Betamethasone valerate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.5 Budesonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.6 Desonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.7 Diflorasone diacetate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.8 Fluticasone propionate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.9 Hydrocortisone buteprate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.10 Mometasone furoate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point	15	2311	Std. Mean Difference (IV, Random, 95% CI)	-0.89 [-1.06, -0.72]
(IAGI/TSS/PASI/PAGI)	1)	2311	Std. Wear Difference (17, Randolli, 7570 Cl)	-0.07 [-1.00, -0.72]
5.1 Betamethasone	3	832	Std. Mean Difference (IV, Random, 95% CI)	-0.80 [-0.96, -0.64]
dipropionate OD	5	032	Std. Mean Difference (1V, Kandolli, 95% CI)	-0.80 [-0.90, -0.04]
5.2 Betamethasone	6	527	Std Mars Difference (IV Denderer 050/ CI)	1 25 [1 5(1 15]
	4	537	Std. Mean Difference (IV, Random, 95% CI)	-1.35 [-1.56, -1.15]
dipropionate twice daily	1	20		0.05 [1.(2, 0.27]
5.3 Betamethasone	1	38	Std. Mean Difference (IV, Random, 95% CI)	-0.95 [-1.62, -0.27]
dipropionate, maintenance	2	0($C = 1 M D^{\circ} C (W D = 1 OS0/C)$	1 22 [1 70 0 00]
5.4 Betamethasone valerate	2	96	Std. Mean Difference (IV, Random, 95% CI)	-1.33 [-1.78, -0.89]
5.5 Budesonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.6 Desonide	1	76	Std. Mean Difference (IV, Random, 95% CI)	-0.81 [-1.34, -0.28]
5.7 Diflorasone diacetate	1	93 292	Std. Mean Difference (IV, Random, 95% CI)	-0.32 [-0.73, 0.09]
5.8 Fluticasone propionate	2	383	Std. Mean Difference (IV, Random, 95% CI)	-0.93 [-1.14, -0.72]
5.9 Hydrocortisone buteprate	1	161	Std. Mean Difference (IV, Random, 95% CI)	-0.46 [-0.77, -0.15]
5.10 Mometasone furoate	1	95	Std. Mean Difference (IV, Random, 95% CI)	-0.75 [-1.17, -0.34]
6 Total withdrawals	9	1673	Risk Difference (M-H, Random, 95% CI)	-0.14 [-0.22, -0.05]
6.1 Betamethasone	2	756	Risk Difference (M-H, Random, 95% CI)	-0.16 [-0.28, -0.04]
dipropionate OD				
6.2 Betamethasone	1	421	Risk Difference (M-H, Random, 95% CI)	-0.06 [-0.12, 0.01]
dipropionate twice daily				
6.3 Betamethasone	2	134	Risk Difference (M-H, Random, 95% CI)	-0.45 [-0.60, -0.30]
dipropionate, maintenance				
6.4 Betamethasone valerate	1	80	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
6.5 Budesonide	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
6.6 Desonide	1	80	Risk Difference (M-H, Random, 95% CI)	-0.18 [-0.38, 0.02]
6.7 Diflorasone diacetate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.8 Fluticasone propionate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.9 Hydrocortisone buteprate	1	180	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.09, 0.10]
6.10 Mometasone furoate	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
7 Withdrawals due to adverse	9	1292	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.05, 0.02]
events				
7.1 Betamethasone	1	633	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.11, -0.02]
dipropionate OD				
7.2 Betamethasone	1	33	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
dipropionate twice daily	-	55		010 [0111, 0111]
7.3 Betamethasone	2	134	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
dipropionate, maintenance	-	т	Tusk Difference (in 11, Randoni, 7970 CI)	5.0 [0.0 I, 0.0 I]
7.4 Betamethasone valerate	1	80	Risk Difference (M-H, Random, 95% CI)	0.08 [-0.02, 0.17]
7.5 Budesonide	1	22	Risk Difference (M-H, Random, 95% CI)	0.08 [-0.16, 0.16]
7.6 Desonide	1	80	Risk Difference (M-H, Random, 95% CI)	-0.1 [-0.24, 0.04]
7.7 Diflorasone diacetate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
	0	v		5.0 [0.0, 0.0]

7.8 Fluticasone propionate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.9 Hydrocortisone buteprate	1	190	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.02, 0.04]
7.10 Mometasone furoate	1	120	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.11, 0.01]
8 Withdrawals due to treatment	6		Risk Difference (M-H, Random, 95% CI)	Subtotals only
failure				,
8.1 Betamethasone	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
dipropionate OD				
8.2 Betamethasone	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
dipropionate twice daily				
8.3 Betamethasone	2	130	Risk Difference (M-H, Random, 95% CI)	-0.46 [-0.61, -0.31]
dipropionate, maintenance				
8.4 Betamethasone valerate	1	80	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
8.5 Budesonide	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
8.6 Desonide	1	80	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
8.7 Diflorasone diacetate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.8 Fluticasone propionate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.9 Hydrocortisone buteprate	1	190	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
8.10 Mometasone furoate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9 Adverse events (local)	10	2117	Risk Difference (M-H, Random, 95% CI)	-0.04 [-0.08,00]
9.1 Betamethasone	2	756	Risk Difference (M-H, Random, 95% CI)	-0.10 [-0.15, -0.04]
dipropionate OD				
9.2 Betamethasone	2	454	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.12, 0.03]
dipropionate twice daily				
9.3 Betamethasone	2	134	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
dipropionate, maintenance				
9.4 Betamethasone valerate	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
9.5 Budesonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.6 Desonide	1	80	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
9.7 Diflorasone diacetate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.8 Fluticasone propionate	1	383	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.05, 0.05]
9.9 Hydrocortisone buteprate	1	190	Risk Difference (M-H, Random, 95% CI)	-0.06 [-0.18, 0.07]
9.10 Mometasone furoate	1	120	Risk Difference (M-H, Random, 95% CI)	-0.10 [-0.23, 0.02]
10 Adverse events (systemic)	4	675	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.01, 0.01]
10.1 Betamethasone	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
dipropionate OD				
10.2 Betamethasone	1	421	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
dipropionate twice daily				
10.3 Betamethasone	2	134	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.07, 0.10]
dipropionate, maintenance				
10.4 Budesonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.5 Desonide	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
10.6 Diflorasone diacetate	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
10.7 Fluticasone propionate	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
10.8 Hydrocortisone	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
buteprate				
10.9 Betamethasone valerate	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
10.10 Mometasone furoate	1	120	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	5	540	Std. Mean Difference (IV, Random, 95% CI)	-1.87 [-2.38, -1.36]
1.1 Clobetasol propionate	4	471	Std. Mean Difference (IV, Random, 95% CI)	-1.89 [-2.53, -1.24]
1.2 Halcinonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.3 Halobetasol	1	69	Std. Mean Difference (IV, Random, 95% CI)	-1.81 [-2.37, -1.24]
2 TSS	3	545	Std. Mean Difference (IV, Random, 95% CI)	-1.35 [-1.80, -0.89]
2.1 Clobetasol propionate	3	545	Std. Mean Difference (IV, Random, 95% CI)	-1.35 [-1.80, -0.89]
2.2 Halcinonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.3 Halobetasol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 PASI	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Clobetasol propionate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Halcinonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.3 Halobetasol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	3	487	Std. Mean Difference (IV, Random, 95% CI)	-1.22 [-1.42, -1.02]
4.1 Clobetasol propionate	1	79	Std. Mean Difference (IV, Random, 95% CI)	-1.01 [-1.55, -0.47]
4.2 Halcinonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Halobetasol	2	408	Std. Mean Difference (IV, Random, 95% CI)	-1.25 [-1.46, -1.04]
5 Combined end point (IAGI/TSS/PASI/PAGI)	10	1493	Std. Mean Difference (IV, Random, 95% CI)	-1.56 [-1.87, -1.26]
5.1 Clobetasol propionate	7	1016	Std. Mean Difference (IV, Random, 95% CI)	-1.65 [-2.10, -1.20]
5.2 Halcinonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.3 Halobetasol	3	477	Std. Mean Difference (IV, Random, 95% CI)	-1.36 [-1.65, -1.07]
6 Total withdrawals	8	1181	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.10, 0.01]
6.1 Clobetasol propionate	7	1037	Risk Difference (M-H, Random, 95% CI)	-0.06 [-0.13, 0.01]
6.2 Halcinonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.3 Halobetasol	1	144	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
7 Withdrawals due to adverse events	10	1601	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.01, 0.01]
7.1 Clobetasol propionate	7	1037	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.01]
7.2 Halcinonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.3 Halobetasol	3	564	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
8 Withdrawals due to treatment	8	1189	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.01]
failure				
8.1 Clobetasol propionate	6	845	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.03, 0.01]
8.2 Halcinonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.3 Halobetasol	2	344	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
9 Adverse events (local)	8	1265	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.02, 0.02]
9.1 Clobetasol propionate	6	845	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.03, 0.03]
9.2 Halcinonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.3 Halobetasol	2	420	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
10 Adverse events (systemic)	6	1056	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.01, 0.01]
10.1 Clobetasol propionate	3	480	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.02, 0.01]
10.2 Halcinonide	1	156	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
10.3 Halobetasol	2	420	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]

Comparison 3. Corticosteroid (very potent) versus placebo

Comparison 4. Dithranol versus placebo

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 TSS	3	94	Std. Mean Difference (IV, Random, 95% CI)	-1.06 [-1.66, -0.46]
3 PASI	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	3	94	Std. Mean Difference (IV, Random, 95% CI)	-1.06 [-1.66, -0.46]
6 Total withdrawals	4	124	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.09, 0.09]
7 Withdrawals due to adverse events	3	104	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
8 Withdrawals due to treatment failure	2	44	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
9 Adverse events (local)	3	94	Risk Difference (M-H, Random, 95% CI)	0.26 [-0.30, 0.82]
10 Adverse events (systemic)	1	20	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.35, 0.35]

Comparison 5. Vitamin D combination products versus placebo

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	5	2264	Std. Mean Difference (IV, Random, 95% CI)	-1.44 [-1.76, -1.12]
1.1 Combination calcipotriol/betamethasone dipropionate, once daily	4	1416	Std. Mean Difference (IV, Random, 95% CI)	-1.21 [-1.50, -0.91]
1.2 Combination calcipotriol/betamethasone dipropionate, twice daily	2	848	Std. Mean Difference (IV, Random, 95% CI)	-1.90 [-2.09, -1.71]
2 TSS	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
2.1 Combination calcipotriol/ betamethasone dipropionate, once daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Combination calcipotriol/ betamethasone dipropionate, twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 PASI	5	2263	Std. Mean Difference (IV, Random, 95% CI)	-1.24 [-1.53, -0.95]
3.1 Combination calcipotriol/betamethasone dipropionate, once daily	4	1414	Std. Mean Difference (IV, Random, 95% CI)	-1.14 [-1.57, -0.70]
3.2 Combination calcipotriol/betamethasone dipropionate, twice daily	2	849	Std. Mean Difference (IV, Random, 95% CI)	-1.41 [-1.86, -0.97]
4 PAGI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected

4.1 Combination calcipotriol/betamethasone	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
dipropionate, once daily 4.2 Combination calcipotriol/ betamethasone dipropionate, twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	5	2264	Std. Mean Difference (IV, Random, 95% CI)	-1.44 [-1.76, -1.12]
5.1 Combination calcipotriol/betamethasone dipropionate, once daily	4	1416	Std. Mean Difference (IV, Random, 95% CI)	-1.21 [-1.50, -0.91]
5.2 Combination calcipotriol/betamethasone dipropionate, twice daily	2	848	Std. Mean Difference (IV, Random, 95% CI)	-1.90 [-2.09, -1.71]
6 Total withdrawals	5	2340	Risk Difference (M-H, Random, 95% CI)	-0.12 [-0.17, -0.07]
6.1 Combination calcipotriol/betamethasone dipropionate, once daily	4	1483	Risk Difference (M-H, Random, 95% CI)	-0.15 [-0.22, -0.09]
6.2 Combination calcipotriol/betamethasone dipropionate, twice daily	2	857	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.12, -0.03]
7 Withdrawals due to adverse events	3	1723	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.11, -0.04]
7.1 Combination calcipotriol/betamethasone dipropionate, once daily	3	1280	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.11, -0.03]
7.2 Combination calcipotriol/betamethasone dipropionate, twice daily	1	443	Risk Difference (M-H, Random, 95% CI)	-0.10 [-0.14, -0.05]
8 Withdrawals due to treatment failure	1	802	Risk Difference (M-H, Random, 95% CI)	-0.09 [-0.12, -0.06]
8.1 Combination calcipotriol/betamethasone dipropionate, once daily	1	359	Risk Difference (M-H, Random, 95% CI)	-0.09 [-0.13, -0.05]
8.2 Combination calcipotriol/betamethasone dipropionate, twice daily	1	443	Risk Difference (M-H, Random, 95% CI)	-0.09 [-0.13, -0.05]
9 Adverse events (local)	5	2334	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.08, -0.02]
9.1 Combination calcipotriol/betamethasone dipropionate, once daily	4	1479	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.11, -0.02]
9.2 Combination calcipotriol/betamethasone dipropionate, twice daily	2	855	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.08, 0.01]
10 Adverse events (systemic)	1	412	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.1 Combination calcipotriol/betamethasone dipropionate, once daily	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Combination calcipotriol/betamethasone dipropionate, twice daily	1	412	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
IAGI	8		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
1.1 Aloe vera extract 0.5% hydrophilic cream, three times per day	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Anti-IL-8 monoclonal antibody cream	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.3 Betamethasone 17-valerate 21-acetate plus tretinoin plus salicylic acid	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.4 Caffeine (topical) 10%, TD	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.5 Calcipotriene 0.005%ointment + nicotinamide 0.05% or 0.1% or 0.7% or 1.4%, twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.6 Dead Sea salts emollient lotion	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.7 Fish oil plus occlusion	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.8 Herbal skin care (Dr Michaels® cleansing gel, ointment and skin conditioner) , twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.9 Hexafluoro-1,25- dihydroxyvitamin D3	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.10 Indigo naturalis 1.4% ointment	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.11 Kukui nut oil, TD	1		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
1.12 <i>Mahonia aquifolium</i> (Reliéva™), twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.13 Methotrexate gel	1		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
1.14 Mycophenolic acid ointment	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.15 NG-monomethyl-L- arginine (L-NMMA) cream	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.16 Nicotinamide 1.4%, twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.17 Oleum horwathiensis (Psoricur®)	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.18 Omega-3- polyunsaturated fatty acids ointment	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.19 Platelet aggregation activating factor (PAF)(Ro 24-0238)	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

1.20 Polymyxin B cream 200, 000 U/g	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.21 PTH (1-34) in	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
Novasome A® liposomal				
cream, twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	
1.22 Sirolimus (topical), 2.2% for 6 wks, then 8% for a	0		Std. Mean Difference (1V, Random, 93% CI)	0.0 [0.0, 0.0]
further 6 wks				
1.23 Tacrolimus ointment	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.24 Tar	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.25 Tazarotene	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.26 Theophylline 1%	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
ointment, twice daily	-			
TSS	17		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 Aloe vera extract 0.5%	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
hydrophilic cream, three times per day				
2.2 Anti-IL-8 monoclonal	1	89	Std. Mean Difference (IV, Random, 95% CI)	-0.70 [-1.13, -0.27]
antibody cream				
2.3 Betamethasone 17-valerate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
21-acetate plus tretinoin plus salicylic acid				
2.4 Caffeine (topical) 10%, TD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.5 Calcipotriene 0.005%	1	192	Std. Mean Difference (IV, Random, 95% CI)	-0.48 [-0.81, -0.15]
ointment + nicotinamide				
0.05% or 0.1% or 0.7% or				
1.4%, twice daily				
2.6 Dead Sea salts emollient	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
lotion				
2.7 Fish oil plus occlusion	1	50	Std. Mean Difference (IV, Random, 95% CI)	-1.05 [-1.64, -0.46]
2.8 Herbal skin care (Dr Michaels® cleansing gel, ointment and skin conditioner) , twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.9 Hexafluoro-1,25-	1	30	Std. Mean Difference (IV, Random, 95% CI)	-1.13 [-1.91, -0.35]
dihydroxyvitamin D3, twice daily	-	50		
2.10 Indigo naturalis 1.4% ointment	2	88	Std. Mean Difference (IV, Random, 95% CI)	-1.64 [-2.13, -1.15]
2.11 Kukui nut oil, TD	1	24	Std. Mean Difference (IV, Random, 95% CI)	0.33 [-0.48, 1.14]
2.12 Mahonia aquifolium	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
(Reliéva™), twice daily				
2.13 Methotrexate gel	1	82	Std. Mean Difference (IV, Random, 95% CI)	-0.48 [-0.92, -0.04]
2.14 Mycophenolic acid ointment	1	14	Std. Mean Difference (IV, Random, 95% CI)	-1.44 [-2.67, -0.22]
2.15	1	34	Std. Mean Difference (IV, Random, 95% CI)	0.08 [-0.60, 0.75]
	1	34	Std. Mean Difference (IV, Random, 95% CI)	0.08 [-0.60, 0.75]

2.17 Oleum horwathiensis	1	42	Std. Mean Difference (IV, Random, 95% CI)	-0.77 [-1.40, -0.14]
(Psoricur®)	0	0		
2.18 Omega-3- polyunsaturated fatty acids	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
ointment				
2.19 Platelet aggregation	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
activating factor (PAF)(Ro 24-	0	0	Std. Ivican Difference (1V, Kandolii, 99% CI)	0.0 [0.0, 0.0]
0238)				
2.20 Polymyxin B cream	1	30	Std. Mean Difference (IV, Random, 95% CI)	0.13 [-0.59, 0.85]
200,000 U/g	1	50	ord. Thean Difference (17, Faindoni, 7776 Cr)	0.15 [0.99, 0.09]
2.21 PTH (1-34) in	1	30	Std. Mean Difference (IV, Random, 95% CI)	-2.31 [-3.26, -1.36]
Novasome A® liposomal		0.0		,,
cream, twice daily				
2.22 Sirolimus (topical), 2.2%	1	44	Std. Mean Difference (IV, Random, 95% CI)	-0.39 [-0.98, 0.21]
for 6 wks, then 8% for a further				
6 wks				
2.23 Tacrolimus ointment	1	47	Std. Mean Difference (IV, Random, 95% CI)	0.06 [-0.52, 0.63]
2.24 Tar	1	36	Std. Mean Difference (IV, Random, 95% CI)	-0.45 [-1.11, 0.22]
2.25 Tazarotene	1	318	Std. Mean Difference (IV, Random, 95% CI)	-0.86 [-1.11, -0.62]
2.26 Theophylline 1%	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
ointment, twice daily				
3 PASI	9		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
3.1 Aloe vera extract 0.5%	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
hydrophilic cream, three times				
per day	0			
3.2 Anti-IL-8 monoclonal	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
antibody cream	1			
3.3 Betamethasone 17-valerate	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
21-acetate plus tretinoin plus salicylic acid				
3.4 Caffeine (topical) 10%,	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
TD	1		std. Wear Difference (17, Randolli, 7970 Cl)	0.0 [0.0, 0.0]
3.5 Calcipotriene 0.005%	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
ointment + nicotinamide 0.	0			0.0 [0.0, 0.0]
05% or 0.1% or 0.7% or 1.				
4%, twice daily				
3.6 Dead Sea salts emollient	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
lotion, 30%				
3.7 Fish oil plus occlusion	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
3.8 Herbal skin care	1		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
(Dr Michaels® cleansing				
gel, ointment and skin				
conditioner), twice daily				
3.9 Hexafluoro-1,25-	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
dihydroxyvitamin D3, twice				
daily	<u>^</u>			
3.10 Indigo naturalis 1.4%	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
ointment	1		Std. Mean Difference (IV, Random, 95% CI)	
3.11 Kukui nut oil, twice daily	1			0.0 [0.0, 0.0]
3.12 <i>Mahonia aquifolium</i> (Reliéva™), twice daily	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
(Reneva), twice daily				

3.13 Methotrexate gel	1	$C_{\rm L} = 1 M D^{1} (\Gamma = 1) O(1 - 1) O(1 - 1)$	
	-	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
3.14 Mycophenolic acid	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
ointment			
3.15 NG-monomethyl-L- arginine (L-NMMA) cream	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.16 Nicotinamide 1.4%, twice daily	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.17 Oleum horwathiensis (Psoricur®)	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.18 Omega-3- polyunsaturated fatty acids ointment	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.19 Platelet aggregation activating factor (PAF)(Ro 24- 0238)	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.20 Polymyxin B cream 200, 000 U/g	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.21 PTH (1-34) in Novasome A® liposomal cream, twice daily	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.22 Sirolimus (topical), 2. 2% for 6 wks, then 8% for a further 6 wks	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.23 Tacrolimus ointment	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
3.24 Tar	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
3.25 Tazarotene	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
3.26 Theophylline 1% ointment, twice daily	1	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	2	Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
4.1 Aloe vera extract 0.5% hydrophilic cream, three times per day	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Anti-IL-8 monoclonal antibody cream	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Betamethasone 17-valerate 21-acetate plus tretinoin plus salicylic acid	1	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.4 Caffeine (topical) 10%, TD	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.5 Calcipotriene 0.005% ointment + nicotinamide 0. 05% or 0.1% or 0.7% or 1. 4%, twice daily	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.6 Dead Sea salts emollient lotion, 30%	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.7 Fish oil plus occlusion	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.8 Herbal skin care (Dr Michaels® cleansing gel, ointment and skin conditioner)	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

4.9 Hexafluoro-1,25- dihydroxyvitamin D3, twice	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
daily				
4.10 Indigo naturalis 1.4% ointment	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.11 Kukui nut oil, TD	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.12 Mahonia aquifolium	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
(Reliéva [™]), twice daily	0		Std. Ivitan Difference (1V, Kandoli, 7770 Cl)	0.0[0.0, 0.0]
4.13 Methotrexate gel	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.14 Mycophenolic acid	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
ointment	0		Stu: Wear Difference (17, Randolli, 7970 Cl)	0.0 [0.0, 0.0]
4.15 NG-monomethyl-L-	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
arginine (L-NMMA) cream	0		Std. Mean Difference (1V, Randolli, 99% CI)	0.0[0.0, 0.0]
4.16 Nicotinamide 1.4%,	0		Std Maan Difference (IV Bandom 05% CI)	[0,0,0,0]
	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0\;[0.0,0.0]$
twice daily	0			0.0[0.0.0]
4.17 Oleum horwathiensis	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
(Psoricur®)				
4.18 Omega-3-	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
polyunsaturated fatty acids				
ointment				
4.19 Platelet aggregation	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
activating factor (PAF)(Ro 24-				
0238)				
4.20 Polymyxin B cream 200,	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
000 U/g				
4.21 PTH (1-34) in	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
Novasome A® liposomal				
cream, twice daily				
4.22 Sirolimus (topical), 2.	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
2% for 6 wks, then 8% for a				
further 6 wks				
4.23 Tacrolimus ointment	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
4.24 Tar	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
4.25 Tazarotene	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
4.26 Theophylline 1%	0		Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
ointment, twice daily				
5 Combined end point	26		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
(IAGI/TSS/PASI/PAGI)				
5.1 Aloe vera extract 0.5%	1	60	Std. Mean Difference (IV, Random, 95% CI)	-1.58 [-2.16, -0.99]
hydrophilic cream, three times				
per day				
5.2 Anti-IL-8 monoclonal	1	89	Std. Mean Difference (IV, Random, 95% CI)	-0.59 [-1.01, -0.16]
antibody cream				
5.3 Betamethasone 17-valerate	1	81	Std. Mean Difference (IV, Random, 95% CI)	-0.76 [-1.21, -0.31]
21-acetate plus tretinoin plus	-	51		5., 6 [1.21, 0.51]
salicylic acid				
5.4 Caffeine (topical) 10%,	1	78	Std. Mean Difference (IV, Random, 95% CI)	-0.39 [-0.84, 0.06]
TD	-	, 0		5.55 [5.61, 6.66]

5.5 Calcipotriene 0.005% ointment + nicotinamide 0.05% or 0.1% or 0.7% or	1	192	Std. Mean Difference (IV, Random, 95% CI)	-0.48 [-0.81, -0.15]
1.4%, twice daily				
5.6 Dead Sea salts emollient lotion, 30%	1	19	Std. Mean Difference (IV, Random, 95% CI)	0.57 [-0.36, 1.51]
5.7 Fish oil plus occlusion	1	50	Std. Mean Difference (IV, Random, 95% CI)	-1.05 [-1.64, -0.46]
5.8 Herbal skin care	1	24	Std. Mean Difference (IV, Random, 95% CI)	-2.96 [-4.19, -1.74]
(Dr Michaels® cleansing gel, ointment and skin				
conditioner), twice daily				
5.9 Hexafluoro-1,25- dihydroxyvitamin D3	1	30	Std. Mean Difference (IV, Random, 95% CI)	-0.62 [-1.35, 0.12]
5.10 Indigo naturalis 1.4% ointment	2	88	Std. Mean Difference (IV, Random, 95% CI)	-2.09 [-2.62, -1.56]
5.11 Kukui nut oil, TD	1	24	Std. Mean Difference (IV, Random, 95% CI)	0.0 [-0.80, 0.80]
5.12 <i>Mahonia aquifolium</i> (Reliéva TM), twice daily	1	200	Std. Mean Difference (IV, Random, 95% CI)	-0.77 [-1.06, -0.48]
5.13 Methotrexate gel	2	142	Std. Mean Difference (IV, Random, 95% CI)	-1.05 [-2.04, -0.06]
5.14 Mycophenolic acid ointment	1	14	Std. Mean Difference (IV, Random, 95% CI)	-1.44 [-2.67, -0.22]
5.15	1	34	Std. Mean Difference (IV, Random, 95% CI)	0.08 [-0.60, 0.75]
NG-monomethyl-L-arginine (L-NMMA) cream				
5.16 Nicotinamide 1.4%, twice daily	1	96	Std. Mean Difference (IV, Random, 95% CI)	-0.20 [-0.60, 0.20]
5.17 Oleum horwathiensis (Psoricur®)	1	42	Std. Mean Difference (IV, Random, 95% CI)	-0.02 [-0.63, 0.58]
5.18 Omega-3- polyunsaturated fatty acids	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
ointment	_			
5.19 Platelet aggregation activating factor (PAF)(Ro 24-0238)	1	80	Std. Mean Difference (IV, Random, 95% CI)	-0.07 [-0.50, 0.37]
5.20 Polymyxin B cream 200,000 U/g	1	30	Std. Mean Difference (IV, Random, 95% CI)	0.13 [-0.59, 0.85]
5.21 PTH (1-34) in Novasome A® liposomal cream, twice daily	1	30	Std. Mean Difference (IV, Random, 95% CI)	-2.31 [-3.26, -1.36]
5.22 Sirolimus (topical), 2.2% for 6 wks, then 8% for a further	1	44	Std. Mean Difference (IV, Random, 95% CI)	-0.39 [-0.98, 0.21]
6 wks	1	47	Std. Mean Difference (IV, Random, 95% CI)	0.06 [0.52 .0.62]
5.23 Tacrolimus ointment 5.24 Tar	1	47 36	Std. Mean Difference (IV, Random, 95% CI) Std. Mean Difference (IV, Random, 95% CI)	0.06 [-0.52, 0.63] -0.45 [-1.11, 0.22]
5.25 Tazarotene	1	318	Std. Mean Difference (IV, Random, 95% CI)	-0.49 [-1.11, 0.22]
5.26 Theophylline 1%	1	22	Std. Mean Difference (IV, Random, 95% CI)	-2.87 [-4.13, -1.62]
ointment, twice daily	-			, [
Total withdrawals	23		Risk Difference (M-H, Random, 95% CI)	Subtotals only
6.1 Aloe vera extract 0.5%	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
hydrophilic cream, three times			, ,	-
per day				

6.2 Anti-IL-8 monoclonal antibody cream	1	96	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.12, 0.08]
6.3 Betamethasone 17-valerate	1	85	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.09, 0.09]
21-acetate plus tretinoin plus salicylic acid	1	09	Nisk Difference (wi-11, Kandoni, 7770 Ci)	-0.00 [-0.09, 0.09]
6.4 Caffeine (topical) 10%, TD	1	78	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
6.5 Calcipotriene 0.005% ointment + nicotinamide 0.05% or 0.1% or 0.7% or 1.4%, twice daily	1	192	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.01, 0.08]
6.6 Dead Sea salts emollient lotion	1	24	Risk Difference (M-H, Random, 95% CI)	0.25 [-0.06, 0.56]
6.7 Fish oil plus occlusion	1	50	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
6.8 Herbal skin care (Dr	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
Michaels® cleansing gel, ointment and skin conditioner) , twice daily				
6.9 Hexafluoro-1,25- dihydroxyvitamin D3	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
6.10 Indigo naturalis 1.4% ointment	2	112	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.15, 0.15]
6.11 Kukui nut oil, TD	1	30	Risk Difference (M-H, Random, 95% CI)	-0.13 [-0.42, 0.15]
6.12 <i>Mahonia aquifolium</i> (Reliéva™), twice daily	1	200	Risk Difference (M-H, Random, 95% CI)	-0.23 [-0.32, -0.14]
6.13 Methotrexate gel	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
6.14 Mycophenolic acid ointment	1	14	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.24, 0.24]
6.15 NG-monomethyl-L-arginine (L-NMMA) cream	1	34	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
6.16 Nicotinamide 1.4%, twice daily	1	96	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.04, 0.08]
6.17 Oleum horwathiensis	1	50	Risk Difference (M-H, Random, 95% CI)	0.16 [-0.04, 0.36]
6.18 Omega-3-polyunsaturated fatty	1	146	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.15, 0.15]
acids ointment 6.19 Platelet aggregation activating factor (PAF)(Ro 24-0238)	1	104	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
6.20 Polymyxin B cream 200,000 U/g	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.24, 0.24]
6.21 PTH (1-34) in Novasome A® liposomal cream, twice daily	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
6.22 Sirolimus (topical)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.23 Tacrolimus ointment	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.24 Tar	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.25 Tazarotene	2	1627	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.01, 0.09]
6.26 Theophylline 1%	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
ointment, twice daily				_

7 Withdrawals due to adverse	19		Risk Difference (M-H, Random, 95% CI)	Subtotals only
events 7.1 Aloe vera extract 0.5%	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
hydrophilic cream, three times per day				
7.2 Anti-IL-8 monoclonal antibody cream	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.3 Betamethasone 17-valerate 21-acetate plus tretinoin plus salicylic acid	1	85	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.07, 0.06]
7.4 Caffeine (topical) 10%, TD	1	78	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
7.5 Calcipotriene 0.005% ointment + nicotinamide 0.05% or 0.1% or 0.7% or 1.4%, twice daily	1	192	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
7.6 Dead Sea salts emollient lotion	1	24	Risk Difference (M-H, Random, 95% CI)	0.08 [-0.18, 0.35]
7.7 Fish oil plus occlusion	1	50	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
7.8 Herbal skin care (Dr Michaels® cleansing gel, ointment and skin conditioner)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
, twice daily				
7.9 Hexafluoro-1,25- dihydroxyvitamin D3	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
7.10 Indigo naturalis 1.4% ointment	2	112	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
7.11 Kukui nut oil, TD	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
7.12 <i>Mahonia aquifolium</i> (Reliéva TM), twice daily	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.13 Methotrexate gel	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
7.14 Mycophenolic acid ointment	1	14	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.24, 0.24]
7.15 NG-monomethyl-L-arginine (L-NMMA) cream	1	34	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
7.16 Nicotinamide 1.4%, twice daily	1	96	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
7.17 Oleum horwathiensis	1	50	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
7.18 Omega-3-polyunsaturated fatty acids ointment	1	146	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
7.19 Platelet aggregation activating factor (PAF)(Ro 24- 0238)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.20 Polymyxin B cream 200, 000 U/g	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.21 PTH (1-34) in Novasome A® liposomal cream, twice daily	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
7.22 Sirolimus (topical)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

7.23 Tacrolimus ointment	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.24 Tar	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.25 Tazarotene	2	1627	Risk Difference (M-H, Random, 95% CI)	0.07 [0.05, 0.10]
7.26 Theophylline 1%	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
ointment, twice daily	10			
8 Withdrawals due to treatment	18		Risk Difference (M-H, Random, 95% CI)	Subtotals only
failure		(0)		
8.1 Aloe vera extract 0.5%	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
hydrophilic cream, three times				
per day	0	0		0.0.[0.0.0.0]
8.2 Anti-IL-8 monoclonal	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
antibody cream		05		
8.3 Betamethasone 17-valerate	1	85	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.04, 0.08]
21-acetate plus tretinoin plus				
salicylic acid		70		
8.4 Caffeine (topical) 10%,	1	78	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
TD		102		
8.5 Calcipotriene 0.005%	1	192	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
ointment + nicotinamide				
0.05% or 0.1% or 0.7% or				
1.4%, twice daily	1	24		0.00[0.12_0.20]
8.6 Dead Sea salts emollient lotion	1	24	Risk Difference (M-H, Random, 95% CI)	0.08 [-0.12, 0.29]
8.7 Fish oil plus occlusion	1	50	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
8.8 Herbal skin care (Dr	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
Michaels® cleansing gel,	0	0	Risk Difference (M-ri, Randolli, 99% CI)	0.0[0.0, 0.0]
ointment and skin conditioner)				
, twice daily				
8.9 Hexafluoro-1,25-	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
dihydroxyvitamin D3	1	50	Risk Difference (W-11, Randolli, 7770 Cl)	0.0 [-0.12, 0.12]
8.10 Indigo naturalis 1.4%	1	28	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.13, 0.13]
ointment	1	20	Risk Difference (W-11, Randolli, 7770 Cl)	0.0 [-0.15, 0.15]
8.11 Kukui nut oil, TD	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
8.12 Mahonia aquifolium	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
(Reliéva TM), twice daily	0	0	Risk Difference (W-11, Randolli, 7770 Cl)	0.0 [0.0, 0.0]
8.13 Methotrexate gel	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
8.14 Mycophenolic acid	1	14	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.24, 0.24]
ointment	1	11	Risk Difference (W 11, Randolli, 7770 Cl)	0.0 [0.2], 0.2]
8.15	1	34	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
NG-monomethyl-L-arginine	1	51	Risk Difference (W 11, Randolli, 7770 Cl)	0.0 [0.11, 0.11]
(L-NMMA) cream				
8.16 Nicotinamide 1.4%,	1	96	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
twice daily	1)0		0.0 [0.0 I, 0.0 I]
8.17 Oleum horwathiensis	1	50	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
8.18	1	146	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
Omega-3-polyunsaturated fatty				[
acids ointment				
8.19 Platelet aggregation	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
activating factor (PAF)(Ro 24-		-		
0238)				

8.20 Polymyxin B cream 200, 000 U/g	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.21 PTH (1-34) in	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
Novasome A® liposomal	1	50		0.0 [0.12, 0.12]
cream, twice daily				
8.22 Sirolimus (topical)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.23 Tacrolimus ointment	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.24 Tar	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.25 Tazarotene	2	1627	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.04, 0.01]
8.26 Theophylline 1%	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
ointment, twice daily	-			
9 Adverse events (local)	21		Risk Difference (M-H, Random, 95% CI)	Subtotals only
9.1 Aloe vera extract 0.5%	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
hydrophilic cream, three times per day	-			
9.2 Anti-IL-8 monoclonal	1	92	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.10, 0.14]
antibody cream				[
9.3 Betamethasone 17-valerate	1	85	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.07, 0.06]
21-acetate plus tretinoin plus salicylic acid	-	0,7		
9.4 Caffeine (topical) 10%,	1	78	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.03, 0.13]
TD				
9.5 Calcipotriene 0.005%	1	192	Risk Difference (M-H, Random, 95% CI)	0.13 [-0.02, 0.27]
ointment + nicotinamide				
0.05% or 0.1% or 0.7% or				
1.4%, twice daily				
9.6 Dead Sea salts emollient	1	24	Risk Difference (M-H, Random, 95% CI)	0.08 [-0.18, 0.35]
lotion				
9.7 Fish oil plus occlusion	1	50	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.06, 0.14]
9.8 Herbal skin care	1	24	Risk Difference (M-H, Random, 95% CI)	-0.09 [-0.44, 0.27]
(Dr Michaels® cleansing				
gel, ointment and skin				
conditioner), twice daily				
9.9 Hexafluoro-1,25-	1	30	Risk Difference (M-H, Random, 95% CI)	0.13 [-0.06, 0.33]
dihydroxyvitamin D3				
9.10 Indigo naturalis 1.4%	2	88	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
ointment				
9.11 Kukui nut oil, TD	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
9.12 Mahonia aquifolium	1	200	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.06, 0.02]
(Reliéva™), twice daily				
9.13 Methotrexate gel	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
9.14 Mycophenolic acid	1	14	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.24, 0.24]
ointment				
9.15	1	34	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
NG-monomethyl-L-arginine				
(L-NMMA) cream				
9.16 Nicotinamide 1.4%,	1	96	Risk Difference (M-H, Random, 95% CI)	0.10 [-0.07, 0.28]
twice daily				
9.17 Oleum horwathiensis	1	50	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.06, 0.14]
				-

9.18	1	146	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.02, 0.05]
Omega-3-polyunsaturated fatty				
acids ointment				
9.19 Platelet aggregation	1	104	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.19, 0.19]
activating factor (PAF)(Ro				
24-0238)				
9.20 Polymyxin B cream 200,	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
000 U/g				
9.21 PTH (1-34) in	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
Novasome A® liposomal				
cream, twice daily				
9.22 Sirolimus (topical)	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
9.23 Tacrolimus ointment	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.24 Tar	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.25 Tazarotene	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.26 Theophylline 1%	1	22	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.16, 0.16]
ointment, twice daily				
10 Adverse events (systemic)	12		Risk Difference (M-H, Random, 95% CI)	Subtotals only
10.1 Aloe vera extract 0.5%	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
hydrophilic cream, three times	Ū	0		010 [010, 010]
per day				
10.2 Anti-IL-8 monoclonal	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
antibody cream	0	0		0.0 [0.0, 0.0]
10.3 Betamethasone	1	85	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
17-valerate 21-acetate plus	1	0)	Risk Difference (191-11, Randolli, 7976 Cl)	0.0 [-0.04, 0.04]
tretinoin plus salicylic acid				
10.4 Caffeine (topical) 10%,	0	0	Did Difference (M.H. Dandem 05% CI)	
TD	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
	0	0	Did Difference (M LL Denders 050/ CL)	
10.5 Calcipotriene 0.005% ointment + nicotinamide 0.	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
05% or 0.1% or 0.7% or 1.				
4%, twice daily				
10.6 Dead Sea salts emollient	0	0	Did Difference (M LL Denders 050/ CL)	
lotion	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
	0	0	Did Difference (M LL Denders 050/ CL)	0.0 [0.0, 0.0]
10.7 Fish oil plus occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	
10.8 Herbal skin care (Dr	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
Michaels® cleansing gel,				
ointment and skin conditioner)				
, twice daily	1	20		0.0[0.12,0.12]
10.9 Hexafluoro-1,25-	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
dihydroxyvitamin D3				
10.10 Indigo naturalis 1.4%	2	88	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
ointment		2		
10.11 Kukui nut oil, TD	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.12 Mahonia aquifolium	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
(Reliéva™), twice daily	2	1//		0.0.[.0.02.0.02]
10.13 Methotrexate gel	2	166	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
10.14 Mycophenolic acid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
ointment				

10.15	1	34	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
NG-monomethyl-L-arginine				
(L-NMMA) cream				
10.16 Nicotinamide 1.4%,	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
twice daily				
10.17 Oleum horwathiensis	1	50	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
10.18 Omega-3-	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
polyunsaturated fatty acids				
ointment				
10.19 Platelet aggregation	1	104	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
activating factor (PAF)(Ro				
24-0238)				
10.20 Polymyxin B cream	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
200,000 U/g				
10.21 PTH (1-34) in	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
Novasome A® liposomal				
cream, twice daily				
10.22 Sirolimus (topical)	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
10.23 Tacrolimus ointment	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
10.24 Tar	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
10.25 Tazarotene	2	414	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.26 Theophylline 1%	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
ointment, twice daily				

Comparison 7. Vitamin D analogues versus corticosteroid (potent)

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Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	8		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol vs.	3	1728	Std. Mean Difference (IV, Random, 95% CI)	0.43 [0.28, 0.58]
betamethasone dipropionate				
1.2 Calcipotriol vs.	1	412	Std. Mean Difference (IV, Random, 95% CI)	-0.02 [-0.21, 0.17]
betamethasone valerate				
1.3 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
desoxymetasone				
1.4 Calcipotriol vs. diflorasone	1	256	Std. Mean Difference (IV, Random, 95% CI)	0.27 [0.02, 0.52]
diacetate				
1.5 Calcipotriol vs.	1	99	Std. Mean Difference (IV, Random, 95% CI)	-0.58 [-0.99, -0.18]
fluocinonide				
1.6 Calcitriol vs.	1	258	Std. Mean Difference (IV, Random, 95% CI)	0.21 [-0.04, 0.45]
betamethasone dipropionate				
1.7 Calcitriol vs.	1	30	Std. Mean Difference (IV, Random, 95% CI)	-0.19 [-0.91, 0.53]
betamethasone valerate				
1.8 Tacalcitol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone valerate				
2 TSS	6		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone dipropionate				

2.2 Calcipotriol vs.	1	684	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.41, -0.11]
betamethasone valerate				
2.3 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
desoxymetasone	_			
2.4 Calcipotriol vs. diflorasone	1	256	Std. Mean Difference (IV, Random, 95% CI)	0.40 [0.15, 0.65]
diacetate		00		
2.5 Calcipotriol vs.	1	89	Std. Mean Difference (IV, Random, 95% CI)	-0.50 [-0.92, -0.07]
fluocinonide	1	250		0.27 [0.02, 0.51]
2.6 Calcitriol vs.	1	258	Std. Mean Difference (IV, Random, 95% CI)	0.27 [0.02, 0.51]
betamethasone dipropionate 2.7 Calcitriol vs.	0	0	Std Mars Differences (IV Denderer 050/ CI)	
betamethasone valerate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.8 Tacalcitol vs.	2	1 / 9	Std. Mean Difference (IV, Random, 95% CI)	0 41 [0 00 0 74]
betamethasone valerate	2	148	Std. Mean Difference (IV, Random, 93% CI)	0.41 [0.09, 0.74]
3 PASI	9		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 Calcipotriol vs.	3	1728	Std. Mean Difference (IV, Random, 95% CI)	0.36 [0.22, 0.51]
betamethasone dipropionate	5	1/20	Std. Wean Difference (17, Kandolii, 7576 Ci)	0.90[0.22, 0.91]
3.2 Calcipotriol vs.	4	1505	Std. Mean Difference (IV, Random, 95% CI)	-0.12 [-0.22, -0.02]
betamethasone valerate	1	1)0)	ote. Mean Difference (17, Nandolli, 7976 Ci)	0.12 [0.22, 0.02]
3.3 Calcipotriol vs.	1	20	Std. Mean Difference (IV, Random, 95% CI)	0.15 [-0.73, 1.02]
desoxymetasone	-	20		0.19 [0., 9, 1.02]
3.4 Calcipotriol vs. diflorasone	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
diacetate				
3.5 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
fluocinonide				
3.6 Calcitriol vs.	1	258	Std. Mean Difference (IV, Random, 95% CI)	0.39 [0.14, 0.63]
betamethasone dipropionate				
3.7 Calcitriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
betamethasone valerate				
3.8 Tacalcitol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone valerate				
4 PAGI	2	1080	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.38, -0.14]
4.1 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
betamethasone dipropionate				
4.2 Calcipotriol vs.	2	1080	Std. Mean Difference (IV, Random, 95% CI)	-0.26 [-0.38, -0.14]
betamethasone valerate				
4.3 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
desoxymetasone				
4.4 Calcipotriol vs. diflorasone	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
diacetate	0	0		
4.5 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
fluocinonide 4.6 Calcitriol vs.	0	0		0.0.[0.0.0.0]
4.6 Calcitriol vs. betamethasone dipropionate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.7 Calcitriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	
4./ Calcitriol vs.	0	0	Stu. Ivican Difference (1v, Kandom, 99% CI)	0.0 [0.0, 0.0]
4.8 Tacalcitol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone valerate	0	U		0.0 [0.0, 0.0]
5 Combined end point	14		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
(IAGI/TSS/PASI/PAGI)				- actorate only

5.1 Calcipotriol vs.	3	1728	Std. Mean Difference (IV, Random, 95% CI)	0.43 [0.28, 0.58]
betamethasone dipropionate				
5.2 Calcipotriol vs.	4	1557	Std. Mean Difference (IV, Random, 95% CI)	-0.12 [-0.26, 0.02]
betamethasone valerate	1	20	Sed Mars Difference (W. Danders 050/ CI)	0 15 [0 72 1 02]
5.3 Calcipotriol vs. desoxymetasone	1	20	Std. Mean Difference (IV, Random, 95% CI)	0.15 [-0.73, 1.02]
5.4 Calcipotriol vs. diflorasone	1	256	Std. Mean Difference (IV, Random, 95% CI)	0.27 [0.02, 0.52]
diacetate	1	290	old. With Difference (17, Nandolli, 7770 Ci)	0.27 [0.02, 0.92]
5.5 Calcipotriol vs.	1	99	Std. Mean Difference (IV, Random, 95% CI)	-0.58 [-0.99, -0.18]
fluocinonide				
5.6 Calcitriol vs.	1	258	Std. Mean Difference (IV, Random, 95% CI)	0.21 [-0.04, 0.45]
betamethasone dipropionate				
5.7 Calcitriol vs.	1	30	Std. Mean Difference (IV, Random, 95% CI)	-0.19 [-0.91, 0.53]
betamethasone valerate				
5.8 Tacalcitol vs.	2	148	Std. Mean Difference (IV, Random, 95% CI)	0.41 [0.09, 0.74]
betamethasone valerate		2005		
6 Total withdrawals	11	3995	Risk Difference (M-H, Random, 95% CI)	0.02 [0.00, 0.03]
6.1 Calcipotriol vs. betamethasone dipropionate	3	1739	Risk Difference (M-H, Random, 95% CI)	0.03 [0.01, 0.06]
6.2 Calcipotriol vs.	3	1520	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.01, 0.04]
betamethasone valerate	5	1)20	Risk Difference (Wi-11, Randolli, 77/0 Cl)	0.01 [-0.01, 0.04]
6.3 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
desoxymetasone				
6.4 Calcipotriol vs. diflorasone	1	268	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
diacetate				
6.5 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
fluocinonide				
6.6 Calcitriol vs.	1	258	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.08, 0.03]
betamethasone dipropionate				
6.7 Calcitriol vs.	1	30	Risk Difference (M-H, Random, 95% CI)	0.07 [-0.10, 0.23]
betamethasone valerate	2	100		0.0[0.11.0.11]
6.8 Tacalcitol vs. betamethasone valerate	2	180	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]
7 Withdrawals due to adverse	9	3058	Risk Difference (M-H, Random, 95% CI)	0.01 [00, 0.01]
events)	5050	Risk Difference (W-11, Randolli, 7970 Cl)	0.01 [00, 0.01]
7.1 Calcipotriol vs.	1	956	Risk Difference (M-H, Random, 95% CI)	0.02 [0.00, 0.04]
betamethasone dipropionate				
7.2 Calcipotriol vs.	3	1520	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.00, 0.01]
betamethasone valerate				
7.3 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
desoxymetasone				
7.4 Calcipotriol vs. diflorasone	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
diacetate				
7.5 Calcipotriol vs. fluocinonide	1	114	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.06, 0.03]
7.6 Calcitriol vs.	1	250	Didy Difference (M.H. Dendern, 0504 CI)	0.01 [0.02 0.02]
betamethasone dipropionate	1	258	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.02, 0.03]
7.7 Calcitriol vs.	1	30	Risk Difference (M-H, Random, 95% CI)	0.07 [-0.10, 0.23]
betamethasone valerate	-	20		
7.8 Tacalcitol vs.	2	180	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
betamethasone valerate				

8 Withdrawals due to treatment failure	5	1500	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.01, 0.01]
8.1 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone dipropionate	0	0	Nusk Difference (WEFF, Randolli, 9970 Cl)	0.0 [0.0, 0.0]
8.2 Calcipotriol vs.	2	1099	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.01, 0.01]
betamethasone valerate	2	1099	Risk Difference (M-11, Randolli, 9970 CI)	-0.00 [-0.01, 0.01]
	0	0		0.0.[0.0.0.0]
8.3 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
desoxymetasone	0	0		
8.4 Calcipotriol vs. diflorasone	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
diacetate				
8.5 Calcipotriol vs.	1	113	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
fluocinonide				
8.6 Calcitriol vs.	1	258	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.03, 0.05]
betamethasone dipropionate				
8.7 Calcitriol vs.	1	30	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.12, 0.12]
betamethasone valerate				
8.8 Tacalcitol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
betamethasone valerate				
9 Adverse events (local)	9	3778	Risk Difference (M-H, Random, 95% CI)	0.07 [0.02, 0.11]
9.1 Calcipotriol vs.	3	1739	Risk Difference (M-H, Random, 95% CI)	0.07 [0.04, 0.09]
betamethasone dipropionate	U	-/0/		, [,,]
9.2 Calcipotriol vs.	3	1516	Risk Difference (M-H, Random, 95% CI)	0.12 [-0.02, 0.26]
betamethasone valerate	5	1)10	Nisk Difference (M-11, Randolli, 9970 Cl)	0.12 [-0.02, 0.20]
9.3 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
-	0	0	Risk Difference (M-H, Randolli, 9)% CI)	0.0[0.0, 0.0]
desoxymetasone	0	0		
9.4 Calcipotriol vs. diflorasone	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
diacetate				
9.5 Calcipotriol vs.	1	113	Risk Difference (M-H, Random, 95% CI)	0.10 [-0.02, 0.22]
fluocinonide				
9.6 Calcitriol vs.	1	258	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.05, 0.06]
betamethasone dipropionate				
9.7 Calcitriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone valerate				
9.8 Tacalcitol vs.	1	152	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.07, 0.04]
betamethasone valerate				
10 Adverse events (systemic)	6	2547	Risk Difference (M-H, Random, 95% CI)	00 [-0.00, 0.00]
10.1 Calcipotriol vs.	1	621	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
betamethasone dipropionate				
10.2 Calcipotriol vs.	3	1516	Risk Difference (M-H, Random, 95% CI)	00 [-0.00, 0.00]
betamethasone valerate	5	1)10	Nusk Difference (WEFF, Randolli, 9970 Cl)	.00 [0.00, 0.00]
10.3 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
desoxymetasone	0	0	Risk Difference (M-11, Randolli, 9970 CI)	0.0[0.0, 0.0]
-	0	0		0.0.[0.0.0.0]
10.4 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
diflorasone diacetate	0	0		
10.5 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
fluocinonide				_
10.6 Calcitriol vs.	1	258	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.04, 0.03]
betamethasone dipropionate				
10.7 Calcitriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
betamethasone valerate				

Comparison 8. Vitamin D analogues versus corticosteroid (very potent)

1

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
1.1 Calcipotriol vs. Clobetasol propionate	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 TSS	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Calcipotriol vs. Clobetasol propionate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 PASI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
3.1 Calcipotriol vs. Clobetasol propionate	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
4.1 Calcipotriol vs. Clobetasol propionate	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	2	82	Std. Mean Difference (IV, Random, 95% CI)	-0.06 [-0.57, 0.44]
5.1 Calcipotriol vs. Clobetasol propionate	2	82	Std. Mean Difference (IV, Random, 95% CI)	-0.06 [-0.57, 0.44]
6 Total withdrawals	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected
6.1 Calcipotriol vs. Clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7 Withdrawals due to adverse events	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected
7.1 Calcipotriol vs. Clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8 Withdrawals due to treatment failure	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected
8.1 Calcipotriol vs. Clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9 Adverse events (local)	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected
9.1 Calcipotriol vs. Clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10 Adverse events (systemic)	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected
10.1 Calcipotriol vs. Clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

Topical treatments for chronic plaque psoriasis (Review)

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Comparison 9. Vitamin D combined with corticosteroid versus cort	ticosteroid
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Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	4		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol +	3	1926	Std. Mean Difference (IV, Random, 95% CI)	-0.40 [-0.52, -0.27]
betamethasone dipropionate vs.				
betamethasone dipropionate				
1.2 Calcipotriol +	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone dipropionate vs. clobetasol propionate				
1.3 Calcipotriol + clobetasol propionate vs. clobetasol	1	65	Std. Mean Difference (IV, Random, 95% CI)	-0.69 [-1.22, -0.15]
propionate				
2 TSS	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
2.1 Calcipotriol + betamethasone dipropionate vs.	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone dipropionate				
2.2 Calcipotriol + betamethasone dipropionate vs. clobetasol propionate	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.3 Calcipotriol + clobetasol propionate vs. clobetasol	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
propionate				
3 PASI	3	1876	Std. Mean Difference (IV, Random, 95% CI)	-0.44 [-0.55, -0.33]
3.1 Calcipotriol + betamethasone dipropionate vs. betamethasone dipropionate	3	1876	Std. Mean Difference (IV, Random, 95% CI)	-0.44 [-0.55, -0.33]
3.2 Calcipotriol +	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone dipropionate vs. clobetasol propionate	Ŭ	Ŭ		0.0 [0.0, 0.0]
3.3 Calcipotriol + clobetasol propionate vs. clobetasol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
propionate	1			T 1 1 1
4 PAGI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
4.1 Calcipotriol + betamethasone dipropionate vs. betamethasone dipropionate	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Calcipotriol + betamethasone dipropionate vs. clobetasol propionate	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Calcipotriol + clobetasol propionate vs. clobetasol propionate	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	5		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 Calcipotriol + betamethasone dipropionate vs. betamethasone dipropionate	3	1926	Std. Mean Difference (IV, Random, 95% CI)	-0.40 [-0.52, -0.27]

5.2 Calcipotriol + betamethasone dipropionate	1	122	Std. Mean Difference (IV, Random, 95% CI)	0.45 [0.09, 0.81]
vs. clobetasol propionate 5.3 Calcipotriol + clobetasol propionate vs. clobetasol	1	65	Std. Mean Difference (IV, Random, 95% CI)	-0.69 [-1.22, -0.15]
propionate 6 Total withdrawals	5	2135	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.01]
6.1 Calcipotriol +	3	1948	Risk Difference (M-H, Random, 95% CI)	00 [-0.03, 0.03]
betamethasone dipropionate vs. betamethasone dipropionate	5	1,10		
6.2 Calcipotriol + betamethasone dipropionate vs. clobetasol propionate	1	122	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
6.3 Calcipotriol + clobetasol propionate vs. clobetasol propionate	1	65	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
7 Withdrawals due to adverse events	3		Risk Difference (M-H, Random, 95% CI)	Totals not selected
7.1 Calcipotriol + betamethasone dipropionate vs. betamethasone dipropionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Calcipotriol + betamethasone dipropionate vs. clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.3 Calcipotriol + clobetasol propionate vs. clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8 Withdrawals due to treatment failure	2		Risk Difference (M-H, Random, 95% CI)	Totals not selected
8.1 Calcipotriol + betamethasone dipropionate vs. betamethasone dipropionate	0		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Calcipotriol + betamethasone dipropionate vs. clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.3 Calcipotriol + clobetasol propionate vs. clobetasol propionate	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9 Adverse events (local)	4		Risk Difference (M-H, Random, 95% CI)	Subtotals only
9.1 Calcipotriol + betamethasone dipropionate vs. betamethasone dipropionate	3	1946	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.00, 0.04]
9.2 Calcipotriol + betamethasone dipropionate vs. clobetasol propionate	1	122	Risk Difference (M-H, Random, 95% CI)	-0.04 [-0.13, 0.06]
9.3 Calcipotriol + clobetasol propionate vs. clobetasol propionate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10 Adverse events (systemic)	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected

1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
0	Risk Difference (M-H Random 95% CI)	0.0 [0.0, 0.0]
0	Risk Difference (MTTI, Randolli, 7776 CI)	0.0 [0.0, 0.0]
0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
	1 0 0	0 Risk Difference (M-H, Random, 95% CI)

Comparison 10. Vitamin D alone or in combination versus dithranol

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	5		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol vs. dithranol	4	994	Std. Mean Difference (IV, Random, 95% CI)	-0.43 [-0.85, -0.01]
1.2 Calcitriol vs. dithranol	1	114	Std. Mean Difference (IV, Random, 95% CI)	0.51 [0.13, 0.88]
1.3 Tacalcitol vs. dithranol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 TSS	4		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 Calcipotriol vs. dithranol	2	210	Std. Mean Difference (IV, Random, 95% CI)	-0.54 [-1.16, 0.08]
2.2 Calcitriol vs. dithranol	1	114	Std. Mean Difference (IV, Random, 95% CI)	0.13 [-0.24, 0.50]
2.3 Tacalcitol vs. dithranol	1	84	Std. Mean Difference (IV, Random, 95% CI)	-0.18 [-0.60, 0.25]
3 PASI	5		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
3.1 Calcipotriol vs. dithranol	3		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Calcitriol vs. dithranol	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.3 Tacalcitol vs. dithranol	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	2		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
4.1 Calcipotriol vs. dithranol	2	544	Std. Mean Difference (IV, Random, 95% CI)	-0.05 [-0.90, 0.80]
4.2 Calcitriol vs. dithranol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Tacalcitol vs. dithranol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	8		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
5.1 Calcipotriol vs. dithranol	6		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Calcitriol vs. dithranol	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.3 Tacalcitol vs. dithranol	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 Total withdrawals	7	615	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.06, 0.01]
6.1 Calcipotriol vs. dithranol	5	417	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.07, 0.04]
6.2 Calcitriol vs. dithranol	1	114	Risk Difference (M-H, Random, 95% CI)	-0.10 [-0.25, 0.06]
6.3 Tacalcitol vs. dithranol	1	84	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.16, 0.11]
7 Withdrawals due to adverse events	7	1265	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.06, -0.00]
7.1 Calcipotriol vs. dithranol	6	1151	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.06, 0.00]
7.2 Calcitriol vs. dithranol	1	114	Risk Difference (M-H, Random, 95% CI)	-0.06 [-0.13, 0.02]
7.3 Tacalcitol vs. dithranol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8 Withdrawals due to treatment	5	788	Risk Difference (M-H, Random, 95% CI)	00 [-0.02, 0.02]
failure)	/00	Non Difference (IVI-11, Nandolli, 7970 CI)	.00 [-0.02, 0.02]
8.1 Calcipotriol vs. dithranol	4	674	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.02, 0.02]
8.2 Calcitriol vs. dithranol	1	114	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.08, 0.04]
8.3 Tacalcitol vs. dithranol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

9 Adverse events (local)	9	1543	Risk Difference (M-H, Random, 95% CI)	-0.32 [-0.43, -0.20]
9.1 Calcipotriol vs. dithranol	7	1345	Risk Difference (M-H, Random, 95% CI)	-0.25 [-0.32, -0.17]
9.2 Calcitriol vs. dithranol	1	114	Risk Difference (M-H, Random, 95% CI)	-0.67 [-0.80, -0.54]
9.3 Tacalcitol vs. dithranol	1	84	Risk Difference (M-H, Random, 95% CI)	-0.36 [-0.52, -0.20]
10 Adverse events (systemic)	4	746	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.01, 0.01]
10.1 Calcipotriol vs. dithranol	2	548	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.01]
10.2 Calcitriol vs. dithranol	1	114	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
10.3 Tacalcitol vs. dithranol	1	84	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]

Comparison 11. Vitamin D alone or in combination versus other vitamin D analogue

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	3		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol vs. calcitriol	1	246	Std. Mean Difference (IV, Random, 95% CI)	0.0 [-0.25, 0.25]
1.2 Calcipotriol vs. tacalcitol	1	226	Std. Mean Difference (IV, Random, 95% CI)	-0.47 [-0.73, -0.21]
1.3 Calcipotriol vs. maxacalcitol	1	52	Std. Mean Difference (IV, Random, 95% CI)	0.43 [-0.12, 0.98]
2 TSS	3	589	Std. Mean Difference (IV, Random, 95% CI)	-0.31 [-0.55, -0.06]
2.1 Calcipotriol vs. calcitriol	1	250	Std. Mean Difference (IV, Random, 95% CI)	-0.32 [-0.57, -0.07]
2.2 Calcipotriol vs. tacalcitol	1	287	Std. Mean Difference (IV, Random, 95% CI)	-0.45 [-0.68, -0.22]
2.3 Calcipotriol vs. maxacalcitol	1	52	Std. Mean Difference (IV, Random, 95% CI)	0.13 [-0.41, 0.68]
3 PASI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
3.1 Calcipotriol vs. calcitriol	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Calcipotriol vs. tacalcitol	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.3 Calcipotriol vs. maxacalcitol	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
4.1 Calcipotriol vs. calcitriol	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Calcipotriol vs. tacalcitol	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Calcipotriol vs. maxacalcitol	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	4	539	Std. Mean Difference (IV, Random, 95% CI)	-0.17 [-0.62, 0.27]
5.1 Calcipotriol vs. calcitriol	2	261	Std. Mean Difference (IV, Random, 95% CI)	-0.41 [-1.46, 0.64]
5.2 Calcipotriol vs. tacalcitol	1	226	Std. Mean Difference (IV, Random, 95% CI)	-0.47 [-0.73, -0.21]
5.3 Calcipotriol vs. maxacalcitol	1	52	Std. Mean Difference (IV, Random, 95% CI)	0.43 [-0.12, 0.98]
6 Total withdrawals	3	334	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.04, 0.08]
6.1 Calcipotriol vs. calcitriol	2	274	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.04, 0.09]
6.2 Calcipotriol vs. tacalcitol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.3 Calcipotriol vs. maxacalcitol	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.17, 0.17]
7 Withdrawals due to adverse events	3	334	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.01, 0.06]
7.1 Calcipotriol vs. calcitriol	2	274	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.01, 0.07]
7.2 Calcipotriol vs. tacalcitol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

7.3 Calcipotriol vs. maxacalcitol	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
8 Withdrawals due to treatment	3	334	Risk Difference (M-H, Random, 95% CI)	00 [-0.02, 0.01]
failure				
8.1 Calcipotriol vs. calcitriol	2	274	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.08, 0.07]
8.2 Calcipotriol vs. tacalcitol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.3 Calcipotriol vs.	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
maxacalcitol				
9 Adverse events (local)	2	537	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.05, 0.12]
9.1 Calcipotriol vs. calcitriol	1	250	Risk Difference (M-H, Random, 95% CI)	0.07 [0.01, 0.14]
9.2 Calcipotriol vs. tacalcitol	1	287	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.09, 0.07]
9.3 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
maxacalcitol				
10 Adverse events (systemic)	3	597	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.1 Calcipotriol vs. calcitriol	1	250	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
10.2 Calcipotriol vs. tacalcitol	1	287	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.3 Calcipotriol vs.	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
maxacalcitol				

Comparison 12. Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	11		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol twice daily vs. calcipotriol OM, BMD ON	1	154	Std. Mean Difference (IV, Random, 95% CI)	0.56 [0.23, 0.88]
1.2 Calcipotriol OD vs. combined calcipotriol + BMD OD	2	1194	Std. Mean Difference (IV, Random, 95% CI)	0.66 [0.31, 1.02]
1.3 Calcipotriol twice daily vs. combined calcipotriol + BMD OD	1	377	Std. Mean Difference (IV, Random, 95% CI)	0.27 [0.06, 0.48]
1.4 Calcipotriol twice daily vs. combined calcipotriol + BMD twice daily	3	1804	Std. Mean Difference (IV, Random, 95% CI)	0.66 [0.40, 0.93]
1.5 Calcipotriol twice daily vs. calcipotriol OM, BMV ON	2	510	Std. Mean Difference (IV, Random, 95% CI)	0.27 [-0.19, 0.74]
1.6 Calcipotriol twice daily vs. calcipotriol OM, clobetasone butyrate ON	1	344	Std. Mean Difference (IV, Random, 95% CI)	0.27 [0.05, 0.48]
1.7 Calcipotriol twice daily vs. calcipotriol twice daily + clobetasol propionate twice daily	1	65	Std. Mean Difference (IV, Random, 95% CI)	0.88 [0.34, 1.42]
1.8 Calcipotriol twice daily vs. calcipotriol OM, diflucortolone valerate ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

1.9 Calcipotriol OD vs. calcipotriol OM, fluocinonide acetonide ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.10 Calcipotriol OD vs. combined calcipotriol + hydrocortisone OD	1	408	Std. Mean Difference (IV, Random, 95% CI)	0.14 [-0.06, 0.33]
1.11 Calcitriol twice daily vs. diflucortolone valerate OM, calcitriol ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.12 Tacalcitol OD vs. combined calcipotriol + BMD OD	1	334	Std. Mean Difference (IV, Random, 95% CI)	0.48 [0.26, 0.70]
2 TSS	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
2.1 Calcipotriol twice daily vs. calcipotriol OM, BMD ON	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Calcipotriol OD vs. combined calcipotriol + BMD OD	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.3 Calcipotriol twice daily vs. combined calcipotriol + BMD OD	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.4 Calcipotriol twice daily vs. combined calcipotriol + BMD twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.5 Calcipotriol twice daily vs. calcipotriol OM, BMV ON	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.6 Calcipotriol twice daily vs. calcipotriol OM, clobetasone butyrate ON	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.7 Calcipotriol twice daily vs. calcipotriol twice daily + clobetasol propionate twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.8 Calcipotriol twice daily vs. calcipotriol OM, diflucortolone valerate ON	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.9 Calcipotriol OD vs. calcipotriol OM, fluocinonide acetonide ON	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.10 Calcipotriol OD vs. combined calcipotriol + hydrocortisone OD	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.11 Calcitriol twice daily vs. diflucortolone valerate OM, calcitriol ON	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.12 Tacalcitol OD vs. combined calcipotriol + BMD OD	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 PASI	16		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 Calcipotriol twice daily vs. calcipotriol OM, BMD ON	1	124	Std. Mean Difference (IV, Random, 95% CI)	0.46 [0.10, 0.82]

3.2 Calcipotriol OD vs. combined calcipotriol + BMD	2	1191	Std. Mean Difference (IV, Random, 95% CI)	0.67 [0.23, 1.11]
OD 3.3 Calcipotriol twice daily vs. combined calcipotriol + BMD	4	1204	Std. Mean Difference (IV, Random, 95% CI)	0.52 [0.38, 0.67]
OD 3.4 Calcipotriol twice daily vs. combined calcipotriol + BMD twice daily	3	1744	Std. Mean Difference (IV, Random, 95% CI)	0.64 [0.46, 0.83]
3.5 Calcipotriol twice daily vs. calcipotriol OM, BMV ON	2	515	Std. Mean Difference (IV, Random, 95% CI)	0.43 [-0.07, 0.93]
3.6 Calcipotriol twice daily vs. calcipotriol OM, clobetasone butyrate ON	1	344	Std. Mean Difference (IV, Random, 95% CI)	0.17 [-0.04, 0.38]
3.7 Calcipotriol twice daily vs. calcipotriol twice daily + clobetasol propionate twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.8 Calcipotriol twice daily vs. calcipotriol OM, diflucortolone valerate ON	1	116	Std. Mean Difference (IV, Random, 95% CI)	0.08 [-0.29, 0.44]
3.9 Calcipotriol OD vs. calcipotriol OM, fluocinonide acetonide ON	1	38	Std. Mean Difference (IV, Random, 95% CI)	0.53 [-0.11, 1.18]
3.10 Calcipotriol OD vs. combined calcipotriol + hydrocortisone OD	1	408	Std. Mean Difference (IV, Random, 95% CI)	0.08 [-0.11, 0.28]
3.11 Calcitriol twice daily vs. diflucortolone valerate OM, calcitriol ON	1	142	Std. Mean Difference (IV, Random, 95% CI)	0.24 [-0.09, 0.57]
3.12 Tacalcitol OD vs. combined calcipotriol + BMD OD	1	334	Std. Mean Difference (IV, Random, 95% CI)	0.47 [0.25, 0.69]
4 PAGI	2	399	Std. Mean Difference (IV, Random, 95% CI)	0.49 [0.29, 0.69]
4.1 Calcipotriol twice daily vs. calcipotriol OM, BMD ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Calcipotriol OD vs. combined calcipotriol + BMD OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Calcipotriol twice daily vs. combined calcipotriol + BMD OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.4 Calcipotriol twice daily vs. combined calcipotriol + BMD twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.5 Calcipotriol twice daily vs. calcipotriol OM, BMV ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.6 Calcipotriol twice daily vs. calcipotriol OM, clobetasone butyrate ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

4.7 Calcipotriol twice daily vs. calcipotriol twice daily + clobetasol propionate twice daily	1	65	Std. Mean Difference (IV, Random, 95% CI)	0.70 [0.16, 1.23]
4.8 Calcipotriol twice daily vs. calcipotriol OM, diflucortolone valerate ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.9 Calcipotriol OD vs. calcipotriol OM, fluocinonide acetonide ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.10 Calcipotriol OD vs. combined calcipotriol + hydrocortisone OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.11 Calcitriol twice daily vs. diflucortolone valerate OM, calcitriol ON	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.12 Tacalcitol OD vs. combined calcipotriol + BMD OD	1	334	Std. Mean Difference (IV, Random, 95% CI)	0.46 [0.24, 0.68]
5 Combined end point (IAGI/TSS/PASI/PAGI)	17		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 Calcipotriol twice daily vs. calcipotriol OM, BMD ON	1	154	Std. Mean Difference (IV, Random, 95% CI)	0.56 [0.23, 0.88]
5.2 Calcipotriol OD vs. combined calcipotriol + BMD OD	2	1194	Std. Mean Difference (IV, Random, 95% CI)	0.66 [0.31, 1.02]
5.3 Calcipotriol twice daily vs. combined calcipotriol + BMD OD	4	1204	Std. Mean Difference (IV, Random, 95% CI)	0.43 [0.20, 0.66]
5.4 Calcipotriol twice daily vs. combined calcipotriol + BMD twice daily	3	1804	Std. Mean Difference (IV, Random, 95% CI)	0.66 [0.40, 0.93]
5.5 Calcipotriol twice daily vs. calcipotriol OM, BMV ON	2	510	Std. Mean Difference (IV, Random, 95% CI)	0.27 [-0.19, 0.74]
5.6 Calcipotriol twice daily vs. calcipotriol OM, clobetasone butyrate ON	1	344	Std. Mean Difference (IV, Random, 95% CI)	0.27 [0.05, 0.48]
5.7 Calcipotriol twice daily vs. calcipotriol twice daily + clobetasol propionate twice daily	1	65	Std. Mean Difference (IV, Random, 95% CI)	0.88 [0.34, 1.42]
5.8 Calcipotriol twice daily vs. calcipotriol OM, diflucortolone valerate ON	1	116	Std. Mean Difference (IV, Random, 95% CI)	0.08 [-0.29, 0.44]
5.9 Calcipotriol OD vs. calcipotriol OM, fluocinonide acetonide ON	1	38	Std. Mean Difference (IV, Random, 95% CI)	0.53 [-0.11, 1.18]
5.10 Calcipotriol OD vs. combined calcipotriol + hydrocortisone OD	1	408	Std. Mean Difference (IV, Random, 95% CI)	0.14 [-0.06, 0.33]

5.11 Calcitriol twice daily vs. diflucortolone valerate OM, calcitriol ON	1	142	Std. Mean Difference (IV, Random, 95% CI)	0.24 [-0.09, 0.57]
5.12 Tacalcitol OD vs. combined calcipotriol + BMD OD	1	334	Std. Mean Difference (IV, Random, 95% CI)	0.48 [0.26, 0.70]
6 Total withdrawals	15	5494	Risk Difference (M-H, Random, 95% CI)	0.03 [0.02, 0.05]
6.1 Talcipotriol vs. calcipotriol and corticosteroid	13	4985	Risk Difference (M-H, Random, 95% CI)	0.03 [0.01, 0.05]
6.2 Calcitriol vs. calcitriol and corticosteroid	1	142	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.08, 0.10]
6.3 Tacalcitol vs. calcipotriol and corticosteroid	1	367	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.01, 0.11]
7 Withdrawals due to adverse events	13	4081	Risk Difference (M-H, Random, 95% CI)	0.02 [0.01, 0.03]
7.1 Calcipotriol vs. calcipotriol and corticosteroid	11	3572	Risk Difference (M-H, Random, 95% CI)	0.02 [0.01, 0.03]
7.2 Calcitriol vs. calcitriol and corticosteroid	1	142	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.02, 0.07]
7.3 Tacalcitol vs. calcipotriol and corticosteroid	1	367	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.02, 0.03]
8 Withdrawals due to treatment failure	7	1925	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.00, 0.02]
8.1 Calcipotriol vs. calcipotriol and corticosteroid	7	1925	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.00, 0.02]
8.2 Calcitriol vs. calcitriol and corticosteroid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.3 Tacalcitol vs. calcipotriol and corticosteroid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9 Adverse events (local)	15	5581	Risk Difference (M-H, Random, 95% CI)	0.06 [0.05, 0.08]
9.1 Calcipotriol vs. calcipotriol and corticosteroid	13	5084	Risk Difference (M-H, Random, 95% CI)	0.06 [0.04, 0.08]
9.2 Calcitriol vs. calcitriol and corticosteroid	1	131	Risk Difference (M-H, Random, 95% CI)	0.10 [0.02, 0.19]
9.3 Tacalcitol vs. calcipotriol and corticosteroid	1	366	Risk Difference (M-H, Random, 95% CI)	0.09 [0.02, 0.15]
10 Adverse events (systemic)	6	2099	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.00, 0.00]
10.1 Calcipotriol vs. calcipotriol and corticosteroid	5	1968	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.00, 0.00]
10.2 Calcitriol vs. calcitriol and corticosteroid	1	131	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
10.3 Tacalcitol vs. calcipotriol and corticosteroid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 13. Vitamin D alone or in combination versus other treatments: complex regimens

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	7		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	1	577	Std. Mean Difference (IV, Random, 95% CI)	-0.12 [-0.29, 0.04]
1.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	585	Std. Mean Difference (IV, Random, 95% CI)	0.13 [-0.04, 0.29]
1.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	1	92	Std. Mean Difference (IV, Random, 95% CI)	0.60 [0.18, 1.02]
1.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy), halometasone (w/e) (2 wks); then calcipotriol twice daily (2wks)	1	76	Std. Mean Difference (IV, Random, 95% CI)	0.41 [-0.05, 0.86]
 1.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12) 	1	125	Std. Mean Difference (IV, Random, 95% CI)	-0.19 [-0.54, 0.16]
1.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	1	759	Std. Mean Difference (IV, Random, 95% CI)	0.27 [0.12, 0.41]

1.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy)+ combined calcipotriol + BMD (w/e) (8 wks)	1	753	Std. Mean Difference (IV, Random, 95% CI)	0.51 [0.37, 0.66]
1.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy)+ combined calcipotriol + BMD (w/e) (8 wks)	1	760	Std. Mean Difference (IV, Random, 95% CI)	0.26 [0.11, 0.40]
1.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	596	Std. Mean Difference (IV, Random, 95% CI)	0.24 [0.08, 0.40]
1.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	493	Std. Mean Difference (IV, Random, 95% CI)	0.54 [0.36, 0.72]
2 TSS	1		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/ dy) & combined calcipotriol + BMD (w/e) (8 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.4 Calcipotriol (6 wks) vs. clobetasol propionate (2wks); then calcipotriol (4 wks)	1	92	Std. Mean Difference (IV, Random, 95% CI)	0.63 [0.21, 1.05]
2.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

2.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
, halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks) 2.7 Calcipotriol ON,	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12)				
2.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks) ; then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 PASI	8		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	1	649	Std. Mean Difference (IV, Random, 95% CI)	-0.04 [-0.19, 0.11]

3.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	1	143	Std. Mean Difference (IV, Random, 95% CI)	0.29 [-0.04, 0.62]
3.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	650	Std. Mean Difference (IV, Random, 95% CI)	0.10 [-0.05, 0.25]
3.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	1	38	Std. Mean Difference (IV, Random, 95% CI)	0.66 [0.01, 1.32]
3.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy), halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks)	1	76	Std. Mean Difference (IV, Random, 95% CI)	1.13 [0.64, 1.62]
3.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12)	1	125	Std. Mean Difference (IV, Random, 95% CI)	-0.27 [-0.62, 0.09]
3.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	1	759	Std. Mean Difference (IV, Random, 95% CI)	0.25 [0.10, 0.39]
3.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	753	Std. Mean Difference (IV, Random, 95% CI)	0.59 [0.45, 0.74]
3.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	760	Std. Mean Difference (IV, Random, 95% CI)	0.30 [0.16, 0.45]

3.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	645	Std. Mean Difference (IV, Random, 95% CI)	0.15 [-0.01, 0.30]
3.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4	1	501	Std. Mean Difference (IV, Random, 95% CI)	0.49 [0.31, 0.67]
wks) 4 PAGI	4		Std. Mean Difference (IV, Random, 95% CI)	Subtotale only
4.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	1	577	Std. Mean Difference (IV, Random, 95% CI)	Subtotals only -0.14 [-0.30, 0.02]
4.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	585	Std. Mean Difference (IV, Random, 95% CI)	0.10 [-0.06, 0.26]
4.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy) , halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	1	759	Std. Mean Difference (IV, Random, 95% CI)	0.28 [0.13, 0.42]

4.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol	1	753	Std. Mean Difference (IV, Random, 95% CI)	0.71 [0.56, 0.85]
 + BMD (w/e) (8 wks) 4.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks) 	1	760	Std. Mean Difference (IV, Random, 95% CI)	0.44 [0.29, 0.58]
4.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	596	Std. Mean Difference (IV, Random, 95% CI)	0.23 [0.07, 0.39]
4.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	493	Std. Mean Difference (IV, Random, 95% CI)	0.54 [0.36, 0.72]
5 Combined end point (IAGI/TSS/PASI/PAGI)	9		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	1	577	Std. Mean Difference (IV, Random, 95% CI)	-0.12 [-0.29, 0.04]
5.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	1	143	Std. Mean Difference (IV, Random, 95% CI)	0.29 [-0.04, 0.62]
5.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	585	Std. Mean Difference (IV, Random, 95% CI)	0.13 [-0.04, 0.29]
5.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	1	92	Std. Mean Difference (IV, Random, 95% CI)	0.60 [0.18, 1.02]
5.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	1	38	Std. Mean Difference (IV, Random, 95% CI)	0.66 [0.01, 1.32]

	5.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy), halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks)	1	76	Std. Mean Difference (IV, Random, 95% CI)	0.41 [-0.05, 0.86]
	5.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12)	1	125	Std. Mean Difference (IV, Random, 95% CI)	-0.19 [-0.54, 0.16]
	5.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	1	759	Std. Mean Difference (IV, Random, 95% CI)	0.27 [0.12, 0.41]
	5.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy)+ combined calcipotriol + BMD (w/e) (8 wks)	1	753	Std. Mean Difference (IV, Random, 95% CI)	0.51 [0.37, 0.66]
	5.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	760	Std. Mean Difference (IV, Random, 95% CI)	0.26 [0.11, 0.40]
	5.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	596	Std. Mean Difference (IV, Random, 95% CI)	0.24 [0.08, 0.40]
	5.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	493	Std. Mean Difference (IV, Random, 95% CI)	0.54 [0.36, 0.72]
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0	Total withdrawals	9	(10	Risk Difference (M-H, Random, 95% CI)	Subtotals only
	6.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	1	649	Risk Difference (M-H, Random, 95% CI)	0.05 [0.00, 0.10]

6.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	1	150	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.09, 0.17]
6.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	649	Risk Difference (M-H, Random, 95% CI)	0.08 [0.03, 0.13]
6.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	1	98	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.09, 0.09]
6.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	1	38	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.10, 0.10]
6.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy), halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks)	1	76	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
6.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12)	1	125	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.12, 0.11]
6.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	1	759	Risk Difference (M-H, Random, 95% CI)	0.08 [0.03, 0.14]
6.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	753	Risk Difference (M-H, Random, 95% CI)	0.11 [0.06, 0.17]
6.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	760	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.01, 0.07]

6.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	644	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.01, 0.07]
6.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	501	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.01, 0.12]
7 Withdrawals due to adverse events	8		Risk Difference (M-H, Random, 95% CI)	Subtotals only
7.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	1	150	Risk Difference (M-H, Random, 95% CI)	-0.04 [-0.09, 0.01]
7.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/ dy) & combined calcipotriol + BMD (w/e) (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	1	98	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
7.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	1	38	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.10, 0.10]
7.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy), halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks)	1	76	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
7.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12)	1	125	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.06, 0.03]

7.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	1	759	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.01, 0.02]
7.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	753	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.02]
7.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	760	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.03, 0.01]
7.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks) ; then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	501	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.01, 0.05]
8 Withdrawals due to treatment failure	6		Risk Difference (M-H, Random, 95% CI)	Subtotals only
8.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	1	150	Risk Difference (M-H, Random, 95% CI)	0.21 [0.10, 0.33]
8.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/ dy) & combined calcipotriol + BMD (w/e) (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	1	98	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]

calciportial OM, Huncinonide acticiportial twice daily (4 wks) 8.6 Calciportial (6 wks) 1 76 Risk Difference (M-H, Random, 95% CI) 0.0 [=0.05, 0.05 ws. halometasone OM, calciportial ON (2 wks); then calciportial on (2 wks); then calciportial twice daily (wds); halometasone (wdc) (2 wks); 1 125 Risk Difference (M-H, Random, 95% CI) 0.03 [=0.04, 0.1 cloberasol propionate OM (2 to 4 vks); then calciportial twice 1 125 Risk Difference (M-H, Random, 95% CI) 0.03 [=0.04, 0.1 Cloberasol propionate OM (2 to 4 Vks); then calciportial twice 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then placebo 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] Numment twice daily (8 wks) 8.9 8.9 8.9 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] NDD (4 wks); then calciportial + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] NDD (4 wks); then calciportial + 0 0 Risk Difference (M-H, Random, 95% CI) <th></th> <th></th> <th></th> <th></th> <th></th>					
8.6 Calciporniol (6 vks) 1 76 Risk Difference (M-H, Random, 95% CI) 0.0 [-0.05, 0.05 vs. halometasone OM, calciportiol Twice daily (wldy), halometasone (vle) (2 vks); there calciportiol Twice daily (2 vks); hence calciportiol Twice daily (vkdy), halometasone (vle) (2 vks); 1 125 Risk Difference (M-H, Random, 95% CI) 0.03 [-0.04, 0.1 clobetasol propionate OM (2 to 4 vks); then calciportiol twice daily (to vk 12) vs. calcitriol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 vks); then calciportiol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 vks); then calciportiol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 vks); then calciportiol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 vks); then calciportiol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 vks); then calciportiol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 vks); then calciportiol + 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 vks); then calciportiol + 0 Risk Difference (M-H, Random, 95% CI)	calcipotriol OM, fluocinonide acetonide ON (2 wks); then	1	38	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.10, 0.10]
8.7 Calcipotriol ON, 1 125 Risk Difference (M-H, Random, 95% CI) 0.03 [-0.04, 0.1 clobetasol propionate OM (2 to 4 why, then calcipotriol twice dily (to wk 12) vs. calcitriol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] Q to 4 wks); then calcipotriol twice 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0]	8.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy), halometasone (w/e) (2 wks); then calcipotriol twice daily (2	1	76	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
8.8 Combined calcipotriol + 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then placebo ointment twice daily (8 wks) 8.9 Combined calcipotriol + 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] <td>8.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol</td> <td>1</td> <td>125</td> <td>Risk Difference (M-H, Random, 95% CI)</td> <td>0.03 [-0.04, 0.10]</td>	8.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol	1	125	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.04, 0.10]
BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks) 8.10 Combined calcipotriol + 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks) 8.11 Combined calcipotriol + BMD (w/e) (8 wks) 8.11 Combined calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks) ; then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks) ; then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks) 8.12 Tacalcitol (8 wks) vs. 1 501 Risk Difference (M-H, Random, 95% CI) 0.05 [0.02, 0.03 combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	8.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.10 Combined calcipotriol0Risk Difference (M-H, Random, 95% CI)0.0 [0.0, 0.0]BMD (4 wks); then calcipotriol0Risk Difference (M-H, Random, 95% CI)0.0 [0.0, 0.0]ointment twice daily (8 wks)vs. combined calcipotriol+BMD (4 wks); then calcipotriol0Risk Difference (M-H, Random, 95% CI)0.0 [0.0, 0.0](w/dy)+ combined calcipotriol0Risk Difference (M-H, Random, 95% CI)0.0 [0.0, 0.0]BMD (8 wks); then calcipotriol0Risk Difference (M-H, Random, 95% CI)0.0 [0.0, 0.0](4 wks) vs. combinedcalcipotriol + BMD (4 wks)+501Risk Difference (M-H, Random, 95% CI)0.05 [0.02, 0.08(w/e) (8 wks)8.12 Tacalcitol (8 wks) vs.1501Risk Difference (M-H, Random, 95% CI)0.05 [0.02, 0.08(4 wks); then calcipotriol (4wks)	BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.11 Combined calcipotriol + 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] BMD (8 wks); then calcipotriol (4 wks) vs. combined 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] (4 wks) vs. combined calcipotriol + BMD (4 wks) 0 0 Risk Difference (M-H, Random, 95% CI) 0.0 [0.0, 0.0] (w/e) (8 wks) 8.12 Tacalcitol (8 wks) vs. 1 501 Risk Difference (M-H, Random, 95% CI) 0.05 [0.02, 0.03 combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	8.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy)+ combined calcipotriol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.12 Tacalcitol (8 wks) vs. 1 501 Risk Difference (M-H, Random, 95% CI) 0.05 [0.02, 0.08 combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	8.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks) ; then calcipotriol (w/dy) & combined calcipotriol + BMD	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
	8.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4	1	501	Risk Difference (M-H, Random, 95% CI)	0.05 [0.02, 0.08]
		8		Risk Difference (M-H, Random, 95% CI)	Subtotals only

9.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	1	649	Risk Difference (M-H, Random, 95% CI)	0.11 [0.06, 0.17]
9.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	649	Risk Difference (M-H, Random, 95% CI)	0.11 [0.05, 0.17]
9.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	1	98	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.08, 0.12]
9.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	1	38	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.14, 0.14]
9.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy), halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks)	1	76	Risk Difference (M-H, Random, 95% CI)	0.26 [0.07, 0.45]
9.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12)	1	125	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.10, 0.06]
9.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	1	752	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.07, 0.02]
9.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	743	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.03, 0.05]

9.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	1	749	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.00, 0.08]
9.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) & combined calcipotriol + BMD (w/e) (8 wks)	1	644	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.05, 0.04]
9.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	501	Risk Difference (M-H, Random, 95% CI)	0.06 [0.01, 0.11]
10 Adverse events (systemic)	0	0	Risk Difference (M-H, Random, 95% CI)	$0.0 \ [0.0, 0.0]$
10.1 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.3 Calcipotriol (12 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/ dy) & combined calcipotriol + BMD (w/e) (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.4 Calcipotriol (6 wks) vs. clobetasol propionate (2 wks); then calcipotriol (4 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.5 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide acetonide ON (2 wks); then calcipotriol twice daily (4 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.6 Calcipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then calcipotriol twice daily (w/dy) , halometasone (w/e) (2 wks); then calcipotriol twice daily (2 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

10.7 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol twice daily (to wk 12) vs. calcitriol ON, clobetasol propionate OM (2 to 4 wks); then calcitriol twice daily (to wk 12)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.8 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.9 Combined calcipotriol + BMD (4 wks); then placebo ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.10 Combined calcipotriol + BMD (4 wks); then calcipotriol ointment twice daily (8 wks) vs. combined calcipotriol + BMD (4 wks) ; then calcipotriol (w/dy) + combined calcipotriol + BMD (w/e) (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (w/ dy) & combined calcipotriol + BMD (w/e) (8 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.12 Tacalcitol (8 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 14. Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks)

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected

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1.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.3 Alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 TSS	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.3 Alternating: combined calcipotriol + BMD (4 wks) ; then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 PASI	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.3 Alternating: combined calcipotriol + BMD (4 wks) ; then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

4.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Alternating: combined calcipotriol + BMD (4 wks) ; then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
5.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5.3 Alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 Total withdrawals	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected
6.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.3 Alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7 Withdrawals due to adverse events	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected

7.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.3 Alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8 Withdrawals due to treatment failure	1	Risk Difference (M-H, Random, 95% CI)	Totals not selected
8.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.3 Alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9 Adverse events (local)	1	Risk Difference (M-H, Random, 95% CI)	Totals not selected
9.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.3 Alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	1	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10 Adverse events (systemic)	0	Risk Difference (M-H, Random, 95% CI)	Subtotals only

10.1 Combined calcipotriol + BMD (52 wks) vs. alternating: combined calcipotriol + BMD (4 wks); then calcipotriol (4 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Combined calcipotriol + BMD (52 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.3 Alternating: combined calcipotriol + BMD (4 wks) ; then calcipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	10		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol vs. coal tar	3	139	Std. Mean Difference (IV, Random, 95% CI)	-0.53 [-1.74, 0.68]
1.2 Calcipotriol vs. coal tar polytherapy	2	209	Std. Mean Difference (IV, Random, 95% CI)	-0.59 [-0.87, -0.31]
1.3 Calcipotriol vs. nicotinamide 1.4%, twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
 1.4 Calcipotriol vs. calcipotriol + nicotinamide 1. 4%, twice daily 	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.5 Calcipotriol vs. corticosteroid + salicylic acid	1	200	Std. Mean Difference (IV, Random, 95% CI)	-0.06 [-0.33, 0.22]
1.6 Calcipotriol vs. propylthiouracil cream	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.7 Calcipotriol vs. tazarotene	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.8 Calcipotriol vs. tacrolimus ointment	1	124	Std. Mean Difference (IV, Random, 95% CI)	-0.22 [-0.60, 0.16]
1.9 Calcipotriol vs. tazarotene gel plus mometasone furoate cream	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.10 Calcipotriol vs. vitamin B12 cream	1	26	Std. Mean Difference (IV, Random, 95% CI)	-0.55 [-1.33, 0.24]
1.11 Head-to-head vitamin D alone or in combination: twice daily vs OD	2	728	Std. Mean Difference (IV, Random, 95% CI)	-0.24 [-0.38, -0.09]
1.12 Head-to-head vitamin D alone or in combination: no occlusion vs. occlusion	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 TSS	7		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only

2.1 Calcipotriol vs. coal tar	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Calcipotriol vs. coal tar	1	132	Std. Mean Difference (IV, Random, 95% CI)	-0.51 [-0.86, -0.16]
polytherapy				
2.3 Calcipotriol vs.	1	96	Std. Mean Difference (IV, Random, 95% CI)	-0.09 [-0.49, 0.31]
nicotinamide 1.4%, twice daily		100		0.40 [0.4 (0.50]
2.4 Calcipotriol vs. calcipotriol+nicotinamide	1	192	Std. Mean Difference (IV, Random, 95% CI)	0.19 [-0.14, 0.52]
(0.05%, 0.1%, 0.7%, or				
(0.09%, 0.1%, 0.7%, 0.7%) of 1.4%), twice daily				
2.5 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
corticosteroid + salicylic acid	0	0		0.0 [0.0, 0.0]
2.6 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
propylthiouracil cream				
2.7 Calcipotriol vs. tacrolimus	2	171	Std. Mean Difference (IV, Random, 95% CI)	-0.35 [-1.51, 0.81]
ointment				
2.8 Calcipotriol vs. tazarotene	1	199	Std. Mean Difference (IV, Random, 95% CI)	-0.05 [-0.33, 0.23]
2.9 Calcipotriol vs. tazarotene	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
gel plus mometasone furoate				
cream				
2.10 Calcipotriol vs. vitamin	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \ [0.0, \ 0.0]$
B12 cream				
2.11 Head-to-head vitamin D	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
alone or in combination: twice				
daily vs OD	2	2/7	$C = 1 M D^{\circ} C (W D = 1 OS0/C)$	0.10[2.04.1.60]
2.12 Head-to-head vitamin D alone or in combination: no	2	247	Std. Mean Difference (IV, Random, 95% CI)	-0.18 [-2.04, 1.68]
occlusion vs. occlusion				
3 PASI	9		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 Calcipotriol vs. coal tar	2	109	Std. Mean Difference (IV, Random, 95% CI)	-0.10 [-1.54, 1.35]
3.2 Calcipotriol vs. coal tar	1	87	Std. Mean Difference (IV, Random, 95% CI)	-0.63 [-1.06, -0.20]
polytherapy				
3.3 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
nicotinamide 1.4%, twice daily				
3.4 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
calcipotriol + nicotinamide 1.				
4%, twice daily				
3.5 Calcipotriol vs.	1	160	Std. Mean Difference (IV, Random, 95% CI)	-0.05 [-0.36, 0.26]
corticosteroid + salicylic acid		27		
3.6 Calcipotriol vs. propylthiouracil cream	1	27	Std. Mean Difference (IV, Random, 95% CI)	-2.24 [-3.23, -1.25]
3.7 Calcipotriol vs. tacrolimus	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
ointment	0	0	Std. Mean Difference (17, Kandolii, 9970 CI)	0.0[0.0, 0.0]
3.8 Calcipotriol vs. tazarotene	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.9 Calcipotriol vs. tazarotene	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
gel plus mometasone furoate				[]
cream				
3.10 Calcipotriol vs. vitamin	1	26	Std. Mean Difference (IV, Random, 95% CI)	-0.01 [-0.78, 0.75]
B12 cream				
3.11 Head-to-head vitamin D	3	989	Std. Mean Difference (IV, Random, 95% CI)	-0.12 [-0.25, 0.00]
alone or in combination: twice				
daily vs OD				

3.12 Head-to-head vitamin D alone or in combination: no	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
occlusion vs. occlusion	<i>,</i>			
4 PAGI	6	5 /	Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
4.1 Calcipotriol vs. coal tar	1	54	Std. Mean Difference (IV, Random, 95% CI)	-1.51 [-2.12, -0.90]
4.2 Calcipotriol vs. coal tar polytherapy	1	87	Std. Mean Difference (IV, Random, 95% CI)	-0.56 [-0.99, -0.13]
4.3 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
nicotinamide 1.4%, twice daily	0	0	Std. Mean Difference (IV, Kandom, 95% CI)	0.0[0.0, 0.0]
4.4 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
calcipotriol + nicotinamide 1.	0	0	Std. Mean Difference (1V, Kandolii, 9970 CI)	0.0 [0.0, 0.0]
4%, twice daily				
4.5 Calcipotriol vs.	1	186	Std. Mean Difference (IV, Random, 95% CI)	-0.49 [-0.79, -0.20]
corticosteroid + salicylic acid	1	100	otd. Wear Difference (17, Randolli, 7576 Ci)	0.19 [0.79, 0.20]
4.6 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
propylthiouracil cream	0	0	ote. Wear Difference (17, Randolli, 7576 Ci)	0.0 [0.0, 0.0]
4.7 Calcipotriol vs. tacrolimus	1	124	Std. Mean Difference (IV, Random, 95% CI)	-0.13 [-0.51, 0.24]
ointment	-	121		0.13 [0.91, 0.21]
4.8 Calcipotriol vs. tazarotene	1	38	Std. Mean Difference (IV, Random, 95% CI)	-0.35 [-0.99, 0.29]
4.9 Calcipotriol vs. tazarotene	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
gel plus mometasone furoate				
cream				
4.10 Calcipotriol vs. vitamin	1	26	Std. Mean Difference (IV, Random, 95% CI)	-0.55 [-1.33, 0.24]
B12 cream				
4.11 Head-to-head vitamin D	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
alone or in combination: twice				
daily vs OD				
4.12 Head-to-head vitamin D	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
alone or in combination: no				
occlusion vs. occlusion				
5 Combined end point	19		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
(IAGI/TSS/PASI/PAGI)	2	120		
5.1 Calcipotriol vs. coal tar	3	139	Std. Mean Difference (IV, Random, 95% CI)	-0.53 [-1.74, 0.68]
5.2 Calcipotriol vs. coal tar	2	209	Std. Mean Difference (IV, Random, 95% CI)	-0.59 [-0.87, -0.31]
polytherapy	1	06		0.00[0.(0.0.21]
5.3 Calcipotriol vs.	1	96	Std. Mean Difference (IV, Random, 95% CI)	-0.09 [-0.49, 0.31]
nicotinamide 1.4%, twice daily 5.4 Calcipotriol vs.	1	192	Std. Mean Difference (IV, Random, 95% CI)	0.19 [-0.14, 0.52]
calcipotriol + nicotinamide	1	192	Std. Mean Difference (IV, Kandom, 95% CI)	0.19 [-0.14, 0.52]
(0.05%, 0.1%, 0.7%, or				
1.4%), twice daily				
5.5 Calcipotriol vs.	2	360	Std. Mean Difference (IV, Random, 95% CI)	-0.05 [-0.26, 0.15]
corticosteroid + salicylic acid	-	500		0.09 [0.20, 0.19]
5.6 Calcipotriol vs.	1	27	Std. Mean Difference (IV, Random, 95% CI)	-2.24 [-3.23, -1.25]
propylthiouracil cream				[[]]]]
5.7 Calcipotriol vs. tacrolimus	2	171	Std. Mean Difference (IV, Random, 95% CI)	-0.55 [-1.28, 0.17]
ointment				
5.8 Calcipotriol vs. tazarotene	2	237	Std. Mean Difference (IV, Random, 95% CI)	-0.10 [-0.35, 0.16]
5.9 Calcipotriol vs. tazarotene	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
gel plus mometasone furoate				
cream				

5.10 Calcipotriol vs. vitamin B12 cream	1	26	Std. Mean Difference (IV, Random, 95% CI)	-0.55 [-1.33, 0.24]
5.11 Head-to-head vitamin D alone or in combination: twice daily vs OD	3	988	Std. Mean Difference (IV, Random, 95% CI)	-0.20 [-0.32, -0.07]
5.12 Head-to-head vitamin D alone or in combination: no occlusion vs. occlusion	2	247	Std. Mean Difference (IV, Random, 95% CI)	-0.18 [-2.04, 1.68]
6 Total withdrawals	15		Risk Difference (M-H, Random, 95% CI)	Subtotals only
6.1 Calcipotriol vs. coal tar	2	120	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.12, 0.08]
6.2 Calcipotriol vs. coal tar polytherapy	2	220	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.10, 0.04]
6.3 Calcipotriol vs. nicotinamide 1.4%, twice daily	1	96	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.05, 0.09]
6.4 Calcipotriol vs. calcipotriol + nicotinamide (0.05%, 0.1%, 0.7%, or 1.4%), twice daily	1	192	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.06, 0.07]
6.5 Calcipotriol vs. corticosteroid + salicylic acid	1	160	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.09, 0.17]
6.6 Calcipotriol vs. propylthiouracil cream	1	28	Risk Difference (M-H, Random, 95% CI)	0.07 [-0.16, 0.30]
6.7 Calcipotriol vs. tacrolimus ointment	1	124	Risk Difference (M-H, Random, 95% CI)	-0.13 [-0.25, -0.01]
6.8 Calcipotriol vs. tazarotene	2	254	Risk Difference (M-H, Random, 95% CI)	-0.04 [-0.10, 0.01]
6.9 Calcipotriol vs. tazarotene	1	120	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.15, 0.08]
gel plus mometasone furoate cream				
6.10 Calcipotriol vs. vitamin B12 cream	1	26	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.14, 0.14]
6.11 Head-to-head vitamin D alone or in combination: twice daily vs OD	3	1001	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.03, 0.05]
6.12 Head-to-head vitamin D alone or in combination: no occlusion vs. occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7 Withdrawals due to adverse events	15		Risk Difference (M-H, Random, 95% CI)	Subtotals only
7.1 Calcipotriol vs. coal tar	2	120	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.04, 0.06]
7.2 Calcipotriol vs. coal tar polytherapy	2	210	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.07, 0.03]
7.3 Calcipotriol vs. nicotinamide 1.4%, twice daily	1	96	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
7.4 Calcipotriol vs. calcipotriol + nicotinamide (0.05%, 0.1%, 0.7%, or 1.4%), twice daily	1	192	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
7.5 Calcipotriol vs. corticosteroid + salicylic acid	1	160	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.00, 0.10]
7.6 Calcipotriol vs. propylthiouracil cream	1	28	Risk Difference (M-H, Random, 95% CI)	0.07 [-0.16, 0.30]

7.7 Calcipotriol vs. tacrolimus ointment	1	124	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.08, 0.11]
7.8 Calcipotriol vs. tazarotene	2	254	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.16, 0.05]
7.9 Calcipotriol vs. tazarotene	1	120	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.10, 0.07]
gel plus mometasone furoate cream	-			
7.10 Calcipotriol vs. vitamin B12 cream	1	26	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.14, 0.14]
7.11 Head-to-head vitamin D alone or in combination: twice daily vs OD	3	998	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.01, 0.02]
7.12 Head-to-head vitamin D alone or in combination: no occlusion vs. occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8 Withdrawals due to treatment failure	12		Risk Difference (M-H, Random, 95% CI)	Subtotals only
8.1 Calcipotriol vs. coal tar	1	60	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
8.2 Calcipotriol vs. coal tar	1	88	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.06, 0.07]
polytherapy				
8.3 Calcipotriol vs. nicotinamide 1.4%, twice daily	1	96	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
8.4 Calcipotriol vs. calcipotriol + nicotinamide (0.05%, 0.1%, 0.7%, or 1.4%), twice daily	1	192	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
8.5 Calcipotriol vs. corticosteroid + salicylic acid	1	160	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.07, 0.02]
8.6 Calcipotriol vs. propylthiouracil cream	1	28	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.25, 0.11]
8.7 Calcipotriol vs. tacrolimus ointment	1	124	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.07, 0.03]
8.8 Calcipotriol vs. tazarotene	1	208	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
8.9 Calcipotriol vs. tazarotene gel plus mometasone furoate cream	1	120	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
8.10 Calcipotriol vs. vitamin B12 cream	1	26	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.14, 0.14]
8.11 Head-to-head vitamin D alone or in combination: twice daily vs OD	3	998	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.01, 0.01]
8.12 Head-to-head vitamin D alone or in combination: no occlusion vs. occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9 Adverse events (local)	11		Risk Difference (M-H, Random, 95% CI)	Subtotals only
9.1 Calcipotriol vs. coal tar	2	120	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.06, 0.10]
9.2 Calcipotriol vs. coal tar polytherapy	1	122	Risk Difference (M-H, Random, 95% CI)	0.06 [-0.09, 0.20]
9.3 Calcipotriol vs. nicotinamide 1.4%, twice daily	1	96	Risk Difference (M-H, Random, 95% CI)	-0.15 [-0.32, 0.03]

9.4 Calcipotriol vs. calcipotriol + nicotinamide (0.05%, 0.1%, 0.7%, or	1	192	Risk Difference (M-H, Random, 95% CI)	-0.17 [-0.30, -0.03]
 1.4%), twice daily 9.5 Calcipotriol vs. corticosteroid + salicylic acid 	1	160	Risk Difference (M-H, Random, 95% CI)	0.09 [0.02, 0.15]
9.6 Calcipotriol vs. propylthiouracil cream	1	28	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.25, 0.11]
9.7 Calcipotriol vs. tacrolimus ointment	1	124	Risk Difference (M-H, Random, 95% CI)	-0.19 [-0.37, -0.01]
9.8 Calcipotriol vs. tazarotene	1	204	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.11, 0.06]
9.9 Calcipotriol vs. tazarotene gel plus mometasone furoate cream	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.10 Calcipotriol vs. vitamin B12 cream	1	26	Risk Difference (M-H, Random, 95% CI)	0.23 [-0.06, 0.52]
9.11 Head-to-head vitamin D alone or in combination: twice daily vs OD	2	731	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.06, 0.05]
9.12 Head-to-head vitamin D alone or in combination: no	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
occlusion vs. occlusion 10 Adverse events (systemic)	0		Disk Difference (MH Bandam 050/ CI)	Subtatala anhu
10.1 Calcipotriol vs. coal tar	8 1	60	Risk Difference (M-H, Random, 95% CI) Risk Difference (M-H, Random, 95% CI)	Subtotals only 0.0 [-0.09, 0.09]
10.2 Calcipotriol vs. coal tar	1	88	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
polytherapy				
10.3 Calcipotriol vs. nicotinamide 1.4%, twice daily	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.4 Calcipotriol vs. calcipotriol + nicotinamide 1. 4%, twice daily	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.5 Calcipotriol vs. corticosteroid + salicylic acid	1	160	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.02, 0.02]
10.6 Calcipotriol vs. propylthiouracil cream	1	28	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.13, 0.13]
10.7 Calcipotriol vs. tacrolimus ointment	1	124	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.04, 0.04]
10.8 Calcipotriol vs. tazarotene	1	183	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.05, 0.03]
10.9 Calcipotriol vs. tazarotene gel plus mometasone furoate cream	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.10 Calcipotriol vs. vitamin B12 cream	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.11 Head-to-head vitamin D alone or in combination: twice daily vs OD	1	264	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.12 Head-to-head vitamin D alone or in combination: no occlusion vs. occlusion	1	38	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.10, 0.10]

Comparison 16. Flexural/facial psoriasis: placebo-controlled trials

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
 1.1 Betamethasone valerate 0. 1%, OD 	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Calcipotriol ointment, OD	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.3 Pimecrolimus cream, 1% OD/twice daily	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.4 Tacrolimus ointment 0. 1%, twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2 TSS	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
2.1 Betamethasone valerate 0. 1%, OD	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Calcipotriol ointment, OD	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.3 Pimecrolimus cream, 1% OD/twice daily	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.4 Tacrolimus ointment 0. 1%, twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3 PASI	1		Std. Mean Difference (IV, Random, 95% CI)	Totals not selected
3.1 Betamethasone valerate 0.1%, OD	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.2 Calcipotriol ointment, OD	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.3 Pimecrolimus cream, 1% OD/twice daily	1		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.4 Tacrolimus ointment 0. 1%, twice daily	0		Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	1		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
4.1 Betamethasone valerate 0. 1%, OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.2 Calcipotriol ointment, OD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Pimecrolimus cream, 1% OD/twice daily	1	47	Std. Mean Difference (IV, Random, 95% CI)	-0.65 [-1.24, -0.06]
4.4 Tacrolimus ointment 0. 1%, twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	2		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 Betamethasone valerate 0.1%, OD	1	36	Std. Mean Difference (IV, Random, 95% CI)	-2.83 [-3.79, -1.88]
5.2 Calcipotriol ointment, OD	1	38	Std. Mean Difference (IV, Random, 95% CI)	-1.08 [-1.77, -0.40]
5.3 Pimecrolimus cream, 1% OD/twice daily	2	86	Std. Mean Difference (IV, Random, 95% CI)	-0.86 [-1.30, -0.41]

5.4 Tacrolimus ointment 0. 1%, twice daily	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
6 Total withdrawals	3		Risk Difference (M-H, Random, 95% CI)	Subtotals only
6.1 Betamethasone valerate 0.1%, OD	1	40	Risk Difference (M-H, Random, 95% CI)	0.10 [-0.08, 0.28]
6.2 Calcipotriol ointment, OD	1	40	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.14, 0.14]
6.3 Pimecrolimus cream, 1% OD/twice daily	2	97	Risk Difference (M-H, Random, 95% CI)	-0.06 [-0.16, 0.04]
6.4 Tacrolimus ointment 0.1%, twice daily	1	167	Risk Difference (M-H, Random, 95% CI)	-0.17 [-0.30, -0.03]
7 Withdrawals due to adverse events	3		Risk Difference (M-H, Random, 95% CI)	Subtotals only
7.1 Betamethasone valerate 0.1%, OD	1	40	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.09, 0.09]
7.2 Calcipotriol ointment, OD	1	40	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.09, 0.09]
7.3 Pimecrolimus cream, 1% OD/twice daily	2	97	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.05, 0.05]
7.4 Tacrolimus ointment 0.1%, twice daily	1	167	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.06, 0.03]
8 Withdrawals due to treatment failure	3		Risk Difference (M-H, Random, 95% CI)	Subtotals only
8.1 Betamethasone valerate 0.1%, OD	1	40	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.18, 0.08]
8.2 Calcipotriol ointment, OD	1	40	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.18, 0.08]
8.3 Pimecrolimus cream, 1% OD/twice daily	2	97	Risk Difference (M-H, Random, 95% CI)	-0.04 [-0.13, 0.04]
8.4 Tacrolimus ointment 0.1%, twice daily	1	167	Risk Difference (M-H, Random, 95% CI)	-0.11 [-0.19, -0.02]
9 Adverse events (local)	3		Risk Difference (M-H, Random, 95% CI)	Subtotals only
9.1 Betamethasone valerate 0.1%, OD	1	40	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.18, 0.08]
9.2 Calcipotriol ointment, OD	1	40	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.11, 0.21]
9.3 Pimecrolimus cream 1%, OD/twice daily	2	97	Risk Difference (M-H, Random, 95% CI)	0.08 [-0.15, 0.31]
9.4 Tacrolimus ointment 0.1%, twice daily	1	167	Risk Difference (M-H, Random, 95% CI)	-0.17 [-0.30, -0.03]
10 Adverse events (systemic)	1		Risk Difference (M-H, Random, 95% CI)	Totals not selected
10.1 Betamethasone valerate 0.1%, OD	0		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Calcipotriol ointment, OD	0		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.3 Pimecrolimus cream 1%, OD/twice daily	0		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.4 Tacrolimus ointment 0.1%, twice daily	1		Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	2		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Calcipotriol vs. BMV	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.2 Calcipotriol vs. calcipotriol + hydrocortisone	1	408	Std. Mean Difference (IV, Random, 95% CI)	0.30 [0.11, 0.50]
1.3 Calcipotriol vs. calcitriol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.4 Calcipotriol vs. pimecrolimus	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.5 Calcitriol vs. tacrolimus	1	49	Std. Mean Difference (IV, Random, 95% CI)	0.42 [-0.15, 0.98]
2 TSS	2		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 Calcipotriol vs. BMV	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
2.2 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
calcipotriol + hydrocortisone				
2.3 Calcipotriol vs. calcitriol	1	150	Std. Mean Difference (IV, Random, 95% CI)	0.61 [0.28, 0.94]
2.4 Calcipotriol vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
pimecrolimus				
2.5 Calcitriol vs. tacrolimus	1	49	Std. Mean Difference (IV, Random, 95% CI)	0.29 [-0.27, 0.85]
3 PASI	2		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 Calcipotriol vs. BMV	1	36	Std. Mean Difference (IV, Random, 95% CI)	2.02 [1.20, 2.84]
3.2 Calcipotriol vs. calcipotriol + hydrocortisone	1	408	Std. Mean Difference (IV, Random, 95% CI)	0.32 [0.12, 0.51]
3.3 Calcipotriol vs. calcitriol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.4 Calcipotriol vs. pimecrolimus	1	39	Std. Mean Difference (IV, Random, 95% CI)	-0.53 [-1.17, 0.11]
3.5 Calcitriol vs. tacrolimus	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4 PAGI	0		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
4.1 Calcipotriol vs. BMV	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0\ [0.0,\ 0.0]$
4.2 Calcipotriol vs. calcipotriol + hydrocortisone	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.3 Calcipotriol vs. calcitriol	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.4 Calcipotriol vs. pimecrolimus	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.5 Calcitriol vs. tacrolimus	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
5 Combined end point (IAGI/TSS/PASI/PAGI)	4	Ŭ	Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 Calcipotriol vs. BMV	1	36	Std. Mean Difference (IV, Random, 95% CI)	2.02 [1.20, 2.84]
5.2 Calcipotriol vs.	1	408	Std. Mean Difference (IV, Random, 95% CI)	0.30 [0.11, 0.50]
calcipotriol + hydrocortisone	1	100	Stal. Intean Difference (17, Randolli, 7976 Ci)	0.50 [0.11, 0.90]
5.3 Calcipotriol vs. calcitriol	1	150	Std. Mean Difference (IV, Random, 95% CI)	0.61 [0.28, 0.94]
5.4 Calcipotriol vs.	1	39	Std. Mean Difference (IV, Random, 95% CI)	-0.53 [-1.17, 0.11]
pimecrolimus				
5.5 Calcitriol vs. tacrolimus	1	49	Std. Mean Difference (IV, Random, 95% CI)	0.42 [-0.15, 0.98]
6 Total withdrawals	4		Risk Difference (M-H, Random, 95% CI)	Subtotals only
6.1 Calcipotriol vs. BMV	1	40	Risk Difference (M-H, Random, 95% CI)	-0.10 [-0.28, 0.08]
6.2 Calcipotriol vs. calcipotriol + hydrocortisone	1	408	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.03, 0.11]
6.3 Calcipotriol vs. calcitriol	1	150	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.11, 0.11]

Comparison 17. Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

6.4 Calcipotriol vs. pimecrolimus	1	40	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.08, 0.18]
6.5 Calcitriol vs. tacrolimus	1	50	Risk Difference (M-H, Random, 95% CI)	0.12 [-0.02, 0.26]
7 Withdrawals due to adverse events	4		Risk Difference (M-H, Random, 95% CI)	Subtotals only
7.1 Calcipotriol vs. BMV	1	40	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.09, 0.09]
7.2 Calcipotriol vs.	1	408	Risk Difference (M-H, Random, 95% CI)	0.06 [0.02, 0.11]
calcipotriol + hydrocortisone 7.3 Calcipotriol vs. calcitriol	1	150	Risk Difference (M-H, Random, 95% CI)	0.00 [0.01 0.19]
7.4 Calcipotriol vs.		40		$0.09 \ [0.01, \ 0.18]$
pimecrolimus	1	40	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.09, 0.09]
7.5 Calcitriol vs. tacrolimus	1	50	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
8 Withdrawals due to treatment	4)0	Risk Difference (M-H, Random, 95% CI)	Subtotals only
failure	4		Kisk Difference (ivi-ri, Kandoni, 95% Ci)	Subtotals only
8.1 Calcipotriol vs. BMV	1	40	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.09, 0.09]
8.2 Calcipotriol vs. calcipotriol + hydrocortisone	1	408	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.01, 0.05]
8.3 Calcipotriol vs. calcitriol	1	150	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
8.4 Calcipotriol vs.	1	40	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.09, 0.09]
pimecrolimus				
8.5 Calcitriol vs. tacrolimus	1	50	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.07, 0.07]
9 Adverse events (local)	3		Risk Difference (M-H, Random, 95% CI)	Subtotals only
9.1 calcipotriol vs. BMV	1	40	Risk Difference (M-H, Random, 95% CI)	0.10 [-0.05, 0.25]
9.2 Calcipotriol vs. calcipotriol + hydrocortisone	1	404	Risk Difference (M-H, Random, 95% CI)	0.15 [0.08, 0.23]
9.3 Calcipotriol vs. calcitriol	1	150	Risk Difference (M-H, Random, 95% CI)	0.09 [0.02, 0.17]
9.4 Calcipotriol vs.	1	40	Risk Difference (M-H, Random, 95% CI)	-0.15 [-0.38, 0.08]
pimecrolimus				
9.5 Calcitriol vs. tacrolimus	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10 Adverse events (systemic)	0		Risk Difference (M-H, Random, 95% CI)	Subtotals only
10.1 Calcipotriol vs. BMV	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.2 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
calcipotriol + hydrocortisone				
10.3 Calcipotriol vs. calcitriol	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.4 Calcipotriol vs.	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
pimecrolimus				
10.5 Calcitriol vs. tacrolimus	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 18. Scalp psoriasis: placebo-controlled trials

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 IAGI	9		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Vitamin D: calcipotriol	2	457	Std. Mean Difference (IV, Random, 95% CI)	-0.72 [-1.28, -0.16]
1.2 Potent steroid:	2	712	Std. Mean Difference (IV, Random, 95% CI)	-1.09 [-1.29, -0.90]
betamethasone dipropionate				
1.3 Potent steroid:	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
betamethasone valerate				

1.4 Very potent steroid:	1	132	Std. Mean Difference (IV, Random, 95% CI)	-1.42 [-1.80, -1.04]
amcinonide	2	450		1 72 [1 00 1 /0]
1.5 Very potent steroid: clobetasol propionate	2	458	Std. Mean Difference (IV, Random, 95% CI)	-1.73 [-1.99, -1.48]
	1	54	Std. Mean Difference (IV, Random, 95% CI)	1 11 [1 60 0 52]
1.6 Very potent steroid: halcinonide	1)4	Std. Mean Difference (IV, Kandom, 99% CI)	-1.11 [-1.69, -0.53]
1.7 Vitamin D in	2	854	Std. Mean Difference (IV, Random, 95% CI)	-0.97 [-1.61, -0.32]
combination: calcipotriol +	Z	0)4	Std. Wear Difference (1V, Kandolli, 9970 CI)	-0.97 [-1.01, -0.32]
BMD				
1.8 Other treatment:	1	20	Std. Mean Difference (IV, Random, 95% CI)	-1.48 [-2.50, -0.47]
betamethasone-17,21-	1	20	Std. Witan Difference (17, Randolli, 7570 Ci)	-1.40 [-2.90, -0.47]
dipropionate plus salicylic acid				
1.9 Other treatment:	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
ciclopirox olamine shampoo	0	0	old. Wear Difference (17, Partioni, 7770 Ci)	0.0 [0.0, 0.0]
1.10 Other treatment:	1	84	Std. Mean Difference (IV, Random, 95% CI)	-1.22 [-1.69, -0.76]
fluocinolone acetonide, plus	1	01	old. Wear Difference (17, Partioni, 7770 Ci)	1.22 [1.0), 0.70]
occlusion				
1.11 Other treatment: salicylic	1	20	Std. Mean Difference (IV, Random, 95% CI)	-0.86 [-1.79, 0.06]
acid				
2 TSS	11		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 Vitamin D: calcipotriol	2	457	Std. Mean Difference (IV, Random, 95% CI)	-0.44 [-0.64, -0.25]
2.2 Potent steroid:	2	712	Std. Mean Difference (IV, Random, 95% CI)	-1.00 [-1.19, -0.81]
betamethasone dipropionate				
2.3 Potent steroid:	1	172	Std. Mean Difference (IV, Random, 95% CI)	-1.40 [-1.75, -1.05]
betamethasone valerate				
2.4 Very potent steroid:	1	126	Std. Mean Difference (IV, Random, 95% CI)	-1.58 [-1.98, -1.18]
amcinonide				
2.5 Very potent steroid:	3	707	Std. Mean Difference (IV, Random, 95% CI)	-1.53 [-1.77, -1.28]
clobetasol propionate				
2.6 Very potent steroid:	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
halcinonide				
2.7 Vitamin D in	2	854	Std. Mean Difference (IV, Random, 95% CI)	-0.92 [-1.42, -0.43]
combination: calcipotriol +				
BMD				
2.8 Other treatment:	1	20	Std. Mean Difference (IV, Random, 95% CI)	-1.15 [-2.11, -0.19]
betamethasone-17,21-				
dipropionate plus salicylic acid				
2.9 Other treatment:	1	37	Std. Mean Difference (IV, Random, 95% CI)	-0.07 [-0.82, 0.68]
ciclopirox olamine shampoo				
2.10 Other treatment:	1	84	Std. Mean Difference (IV, Random, 95% CI)	-0.89 [-1.34, -0.44]
fluocinolone acetonide, plus				
occlusion				
2.11 Other treatment: salicylic	1	20	Std. Mean Difference (IV, Random, 95% CI)	-0.57 [-1.47, 0.32]
acid	0			C 1 1 1
3 PASI	0	0	Std. Mean Difference (IV, Random, 95% CI)	Subtotals only 0.0 [0.0, 0.0]
3.1 Vitamin D: calcipotriol	0	0	Std. Mean Difference (IV, Random, 95% CI)	
3.2 Potent steroid:	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
betamethasone dipropionate 3.3 Potent steroid:	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0.[0.0.0.0]
5.5 Potent steroid: betamethasone valerate	U	0	Std. Mean Difference (1V, Kandom, 93% CI)	0.0 [0.0, 0.0]
betamethasone valerate				

3.4 Very potent steroid: amcinonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.5 Very potent steroid:	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
clobetasol propionate				[,]
3.6 Very potent steroid: halcinonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.7 Vitamin D in combination: calcipotriol + BMD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.8 Other treatment: betamethasone-17,21- dipropionate plus salicylic acid	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.9 Other treatment: ciclopirox olamine shampoo	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.10 Other treatment: fluocinolone acetonide, plus occlusion	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
3.11 Other treatment: salicylic acid	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
PAGI	5		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
4.1 Vitamin D: calcipotriol	2	450	Std. Mean Difference (IV, Random, 95% CI)	-0.66 [-1.28, -0.05]
4.2 Potent steroid: betamethasone dipropionate	1	685	Std. Mean Difference (IV, Random, 95% CI)	-1.23 [-1.43, -1.03]
4.3 Potent steroid: betamethasone valerate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.4 Very potent steroid: amcinonide	1	132	Std. Mean Difference (IV, Random, 95% CI)	-0.97 [-1.33, -0.61]
4.5 Very potent steroid: clobetasol propionate	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.6 Very potent steroid: halcinonide	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.7 Vitamin D in combination: calcipotriol + BMD	2	841	Std. Mean Difference (IV, Random, 95% CI)	-1.00 [-1.79, -0.22]
4.8 Other treatment: betamethasone-17,21- dipropionate plus salicylic acid	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.9 Other treatment: ciclopirox olamine shampoo	1	37	Std. Mean Difference (IV, Random, 95% CI)	-0.11 [-0.86, 0.64]
4.10 Other treatment: fluocinolone acetonide, plus occlusion	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
4.11 Other treatment: salicylic acid	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
Combined end point (IAGI/TSS/PASI/PAGI)	13		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
5.1 Vitamin D: calcipotriol	2	457	Std. Mean Difference (IV, Random, 95% CI)	-0.72 [-1.28, -0.16]
5.2 Potent steroid:	2	712	Std. Mean Difference (IV, Random, 95% CI)	-1.09 [-1.29, -0.90]
betamethasone dipropionate				
5.3 Potent steroid: betamethasone valerate	1	172	Std. Mean Difference (IV, Random, 95% CI)	-1.40 [-1.75, -1.05]

5.4 Very potent steroid: amcinonide	1	132	Std. Mean Difference (IV, Random, 95% CI)	-1.42 [-1.80, -1.04]
5.5 Very potent steroid: clobetasol propionate	4	788	Std. Mean Difference (IV, Random, 95% CI)	-1.57 [-1.81, -1.34]
5.6 Very potent steroid: halcinonide	1	54	Std. Mean Difference (IV, Random, 95% CI)	-1.11 [-1.69, -0.53]
5.7 Vitamin D in combination: calcipotriol + BMD	2	854	Std. Mean Difference (IV, Random, 95% CI)	-0.97 [-1.61, -0.32]
5.8 Other treatment: betamethasone-17,21- dipropionate plus salicylic acid	1	20	Std. Mean Difference (IV, Random, 95% CI)	-1.48 [-2.50, -0.47]
5.9 Other treatment: ciclopirox olamine shampoo	1	37	Std. Mean Difference (IV, Random, 95% CI)	-0.07 [-0.82, 0.68]
5.10 Other treatment: fluocinolone acetonide, plus occlusion	1	84	Std. Mean Difference (IV, Random, 95% CI)	-1.22 [-1.69, -0.76]
5.11 Other treatment: salicylic acid	1	20	Std. Mean Difference (IV, Random, 95% CI)	-0.86 [-1.79, 0.06]
6 Total withdrawals	13		Risk Difference (M-H, Random, 95% CI)	Subtotals only
6.1 Vitamin D: calcipotriol	3	517	Risk Difference (M-H, Random, 95% CI)	-0.02 [-0.08, 0.05]
6.2 Potent steroid: betamethasone dipropionate	1	692	Risk Difference (M-H, Random, 95% CI)	-0.14 [-0.21, -0.06]
6.3 Potent steroid: betamethasone valerate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.4 Very potent steroid: amcinonide	1	165	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.11, 0.11]
6.5 Very potent steroid: clobetasol propionate	5	1006	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.10, 0.04]
6.6 Very potent steroid: halcinonide	1	58	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.13, 0.13]
6.7 Vitamin D in combination: calcipotriol + BMD	2	854	Risk Difference (M-H, Random, 95% CI)	-0.09 [-0.16, -0.03]
6.8 Other treatment: betamethasone-17,21- dipropionate plus salicylic acid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
6.9 Other treatment: ciclopirox olamine shampoo	1	40	Risk Difference (M-H, Random, 95% CI)	-0.15 [-0.38, 0.09]
6.10 Other treatment: fluocinolone acetonide, plus occlusion	1	89	Risk Difference (M-H, Random, 95% CI)	-0.04 [-0.13, 0.04]
6.11 Other treatment: salicylic acid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7 Withdrawals due to adverse events	13		Risk Difference (M-H, Random, 95% CI)	Subtotals only
7.1 Vitamin D: calcipotriol	3	517	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.02, 0.05]
7.2 Potent steroid:	2	712	Risk Difference (M-H, Random, 95% CI)	-0.04 [-0.08, -0.00]
betamethasone dipropionate 7.3 Potent steroid:	1	172	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]
betamethasone valerate				

7.4 Very potent steroid: amcinonide	1	165	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.02, 0.04]
7.5 Very potent steroid:	5	1006	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.01]
clobetasol propionate	-			
7.6 Very potent steroid: halcinonide	1	58	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.06, 0.06]
7.7 Vitamin D in combination: calcipotriol + BMD	1	677	Risk Difference (M-H, Random, 95% CI)	-0.04 [-0.08, 0.00]
7.8 Other treatment: betamethasone-17,21- dipropionate plus salicylic acid	1	20	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.17, 0.17]
7.9 Other treatment: ciclopirox olamine shampoo	1	40	Risk Difference (M-H, Random, 95% CI)	-0.18 [-0.42, 0.05]
7.10 Other treatment: fluocinolone acetonide, plus occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
7.11 Other treatment: salicylic acid	1	20	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.17, 0.17]
8 Withdrawals due to treatment failure	9		Risk Difference (M-H, Random, 95% CI)	Subtotals only
8.1 Vitamin D: calcipotriol	2	457	Risk Difference (M-H, Random, 95% CI)	-0.05 [-0.11, 0.00]
8.2 Potent steroid: betamethasone dipropionate	1	692	Risk Difference (M-H, Random, 95% CI)	-0.10 [-0.16, -0.05]
8.3 Potent steroid: betamethasone valerate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.4 Very potent steroid: amcinonide	1	165	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.05, 0.02]
8.5 Very potent steroid: clobetasol propionate	5	1006	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.05, 0.02]
8.6 Very potent steroid: halcinonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.7 Vitamin D in combination: calcipotriol + BMD	1	677	Risk Difference (M-H, Random, 95% CI)	-0.11 [-0.17, -0.06]
8.8 Other treatment: betamethasone-17,21- dipropionate plus salicylic acid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.9 Other treatment: ciclopirox olamine shampoo	1	40	Risk Difference (M-H, Random, 95% CI)	-0.09 [-0.28, 0.10]
8.10 Other treatment: fluocinolone acetonide, plus occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
8.11 Other treatment: salicylic acid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9 Adverse events (local)	12		Risk Difference (M-H, Random, 95% CI)	Subtotals only
9.1 Vitamin D: calcipotriol	3	510	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.05, 0.04]
9.2 Potent steroid:	2	703	Risk Difference (M-H, Random, 95% CI)	-0.07 [-0.13, -0.01]
betamethasone dipropionate				-
9.3 Potent steroid:	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
betamethasone valerate				

9.4 Very potent steroid: amcinonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9.5 Very potent steroid: clobetasol propionate	4	817	Risk Difference (M-H, Random, 95% CI)	0.00 [-0.03, 0.04]
9.6 Very potent steroid: halcinonide	1	58	Risk Difference (M-H, Random, 95% CI)	-0.03 [-0.12, 0.06]
9.7 Vitamin D in combination: calcipotriol + BMD	2	831	Risk Difference (M-H, Random, 95% CI)	-0.06 [-0.13, 0.02]
9.8 Other treatment: betamethasone-17,21- dipropionate plus salicylic acid	1	20	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.17, 0.17]
9.9 Other treatment: ciclopirox olamine shampoo	1	40	Risk Difference (M-H, Random, 95% CI)	-0.06 [-0.24, 0.13]
9.10 Other treatment: fluocinolone acetonide, plus occlusion	1	89	Risk Difference (M-H, Random, 95% CI)	0.02 [-0.04, 0.08]
9.11 Other treatment: salicylic acid	1	20	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.17, 0.17]
10 Adverse events (systemic)	4		Risk Difference (M-H, Random, 95% CI)	Subtotals only
10.1 Vitamin D: calcipotriol	1	408	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.2 Potent steroid: betamethasone dipropionate	1	692	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.3 Potent steroid: betamethasone valerate	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.4 Very potent steroid: amcinonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.5 Very potent steroid: clobetasol propionate	2	385	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.03, 0.02]
10.6 Very potent steroid: halcinonide	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.7 Vitamin D in combination: calcipotriol + BMD	2	843	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.8 Other treatment: betamethasone-17,21- dipropionate plus salicylic acid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.9 Other treatment: ciclopirox olamine shampoo	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.10 Other treatment: fluocinolone acetonide, plus occlusion	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
10.11 Other treatment: salicylic acid	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]

Comparison 19. Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome or subgroup title	No. of No. of studies participants		Statistical method	Effect size
1 IAGI	9		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
1.1 Vitamin D vs.	2	1676	Std. Mean Difference (IV, Random, 95% CI)	0.48 [0.32, 0.64]
corticosteroid (potent): calcipotriol vs. BMD				
1.2 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMV	2	510	Std. Mean Difference (IV, Random, 95% CI)	0.37 [0.20, 0.55]
 1.3 Vitamin D vs. corticosteroid (very potent) : calcipotriol vs. clobetasol propionate 	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
1.4 Vitamin D + corticosteroid vs. corticosteroid: calcipotriol + BMD vs. BMD	3	2444	Std. Mean Difference (IV, Random, 95% CI)	-0.18 [-0.26, -0.10]
1.5 Vitamin D vs. vitamin D + corticosteroid: calcipotriol vs. calcipotriol + BMD	4	2581	Std. Mean Difference (IV, Random, 95% CI)	0.64 [0.44, 0.84]
1.6 Vitamin D vs. other treatments: calcipotriol vs. coal tar polytherapy	2	748	Std. Mean Difference (IV, Random, 95% CI)	-0.24 [-0.73, 0.25]
2 TSS	10		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
2.1 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMD	2	1676	Std. Mean Difference (IV, Random, 95% CI)	0.45 [0.28, 0.63]
2.2 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMV	2	487	Std. Mean Difference (IV, Random, 95% CI)	0.09 [-0.09, 0.27]
2.3 Vitamin D vs. corticosteroid (very potent): calcipotriol vs. clobetasol propionate	1	151	Std. Mean Difference (IV, Random, 95% CI)	0.37 [0.05, 0.69]
2.4 Vitamin D + corticosteroid vs. corticosteroid: calcipotriol + BMD vs. BMD	3	2444	Std. Mean Difference (IV, Random, 95% CI)	-0.19 [-0.27, -0.11]
2.5 Vitamin D vs. vitamin D + corticosteroid: calcipotriol vs. calcipotriol + BMD	3	1978	Std. Mean Difference (IV, Random, 95% CI)	0.70 [0.56, 0.84]
2.6 Vitamin D vs. other treatments: calcipotriol vs. coal tar polytherapy	3	925	Std. Mean Difference (IV, Random, 95% CI)	-0.30 [-0.84, 0.24]
3 PASI	0		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
3.1 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMD	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]

3.2 Vitamin D vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0 0 [0 0 0 0]
corticosteroid (potent):	0	0	Std. Mean Difference (1V, Kandolii, 95% CI)	0.0 [0.0, 0.0]
calcipotriol vs. BMV				
3.3 Vitamin D vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
corticosteroid (very potent)				
: calcipotriol vs. clobetasol propionate				
3.4 Vitamin D + corticosteroid	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
vs. corticosteroid: calcipotriol +	Ū	Ŭ		0.0 [0.0, 0.0]
BMD vs. BMD				
3.5 Vitamin D vs. vitamin D	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
+ corticosteroid: calcipotriol vs.				
calcipotriol + BMD 3.6 Vitamin D vs. other	0	0	Sed Man Differences (IV Decidence 050/ CI)	0 0 [0 0 0 0]
5.6 Vitamin D vs. other treatments: calcipotriol vs. coal	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
tar polytherapy				
4 PAGI	5		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
4.1 Vitamin D vs.	2	1654	Std. Mean Difference (IV, Random, 95% CI)	0.56 [0.31, 0.81]
corticosteroid (potent):				
calcipotriol vs. BMD	1	460		0 (1 [0 22 0 50]
4.2 Vitamin D vs. corticosteroid (potent):	1	468	Std. Mean Difference (IV, Random, 95% CI)	0.41 [0.22, 0.59]
calcipotriol vs. BMV				
4.3 Vitamin D vs.	0	0	Std. Mean Difference (IV, Random, 95% CI)	0.0 [0.0, 0.0]
corticosteroid (very potent)				
: calcipotriol vs. clobetasol				
propionate	2	2/1/		0.17[0.26_0.00]
4.4 Vitamin D + corticosteroid vs. corticosteroid: calcipotriol +	3	2414	Std. Mean Difference (IV, Random, 95% CI)	-0.17 [-0.25, -0.09]
BMD vs. BMD				
4.5 Vitamin D vs. vitamin D	3	1952	Std. Mean Difference (IV, Random, 95% CI)	0.84 [0.61, 1.08]
+ corticosteroid: calcipotriol vs.				
calcipotriol + BMD				
4.6 Vitamin D vs. other	0	0	Std. Mean Difference (IV, Random, 95% CI)	$0.0 \; [0.0, 0.0]$
treatments: calcipotriol vs. coal tar polytherapy				
5 Combined end point	11		Std. Mean Difference (IV, Random, 95% CI)	Subtotals only
(IAGI/TSS/PASI/PAGI)				,
5.1 Vitamin D vs.	2	1676	Std. Mean Difference (IV, Random, 95% CI)	0.48 [0.32, 0.64]
corticosteroid (potent):				
calcipotriol vs. BMD	2	510		
5.2 Vitamin D vs. corticosteroid (potent):	2	510	Std. Mean Difference (IV, Random, 95% CI)	0.37 [0.20, 0.55]
calcipotriol vs. BMV				
5.3 Vitamin D vs.	1	151	Std. Mean Difference (IV, Random, 95% CI)	0.37 [0.05, 0.69]
corticosteroid (very potent):				
calcipotriol vs. clobetasol				
propionate	2	2666	Sed Moon Difference (W. Day Jaw 050/ CI)	0.10[0.26_0.10]
5.4 Vitamin D + corticosteroid vs. corticosteroid: calcipotriol +	3	2444	Std. Mean Difference (IV, Random, 95% CI)	-0.18 [-0.26, -0.10]
BMD vs. BMD				

5.5 Vitamin D vs. vitamin D + corticosteroid: calcipotriol vs. calcipotriol + BMD	4	2581	Std. Mean Difference (IV, Random, 95% CI)	0.64 [0.44, 0.84]
5.6 Vitamin D vs. other treatments: calcipotriol vs. coal tar polytherapy	3	835	Std. Mean Difference (IV, Random, 95% CI)	-0.45 [-0.92, 0.02]
6 Total withdrawals	10		Risk Difference (M-H, Random, 95% CI)	Subtotals only
6.1 Vitamin D vs.	2	1676	Risk Difference (M-H, Random, 95% CI)	0.07 [-0.04, 0.18]
corticosteroid (potent): calcipotriol vs. BMD				
6.2 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMV	2	516	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.00, 0.08]
6.3 Vitamin D vs. corticosteroid (very potent): calcipotriol vs. clobetasol propionate	2	194	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.07, 0.18]
6.4 Vitamin D + corticosteroid vs. corticosteroid: calcipotriol + BMD vs. BMD	3	2444	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.04, 0.06]
6.5 Vitamin D vs. vitamin D + corticosteroid: calcipotriol vs. calcipotriol + BMD	4	2847	Risk Difference (M-H, Random, 95% CI)	0.11 [0.05, 0.18]
6.6 Vitamin D vs. other treatments: calcipotriol vs. coal tar polytherapy	1	475	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.07, 0.09]
7 Withdrawals due to adverse	10		Risk Difference (M-H, Random, 95% CI)	Subtotals only
events	10			Subtotals only
7.1 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMD	2	1676	Risk Difference (M-H, Random, 95% CI)	0.04 [-0.01, 0.09]
7.2 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMV	2	516	Risk Difference (M-H, Random, 95% CI)	0.03 [0.01, 0.06]
7.3 Vitamin D vs. corticosteroid (very potent): calcipotriol vs. clobetasol propionate	2	194	Risk Difference (M-H, Random, 95% CI)	0.05 [-0.05, 0.15]
7.4 Vitamin D + corticosteroid vs. corticosteroid: calcipotriol + BMD vs. BMD	3	2444	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.01, 0.01]
7.5 Vitamin D vs. vitamin D + corticosteroid: calcipotriol vs. calcipotriol + BMD	4	2847	Risk Difference (M-H, Random, 95% CI)	0.06 [0.02, 0.09]
7.6 Vitamin D vs. other treatments: calcipotriol vs. coal tar polytherapy	1	445	Risk Difference (M-H, Random, 95% CI)	0.08 [0.02, 0.14]
8 Withdrawals due to treatment failure	8		Risk Difference (M-H, Random, 95% CI)	Subtotals only

8.1 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMD	2	1676	Risk Difference (M-H, Random, 95% CI)	0.03 [-0.01, 0.07]
8.2 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMV	2	516	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.01, 0.03]
8.3 Vitamin D vs. corticosteroid (very potent): calcipotriol vs. clobetasol propionate	2	194	Risk Difference (M-H, Random, 95% CI)	0.01 [-0.02, 0.04]
8.4 Vitamin D + corticosteroid vs. corticosteroid: calcipotriol + BMD vs. BMD	3	2444	Risk Difference (M-H, Random, 95% CI)	-0.01 [-0.02, -0.00]
8.5 Vitamin D vs. vitamin D + corticosteroid: calcipotriol vs. calcipotriol + BMD	3	2535	Risk Difference (M-H, Random, 95% CI)	0.05 [0.01, 0.10]
8.6 Vitamin D vs. other treatments: calcipotriol vs. coal tar polytherapy	0	0	Risk Difference (M-H, Random, 95% CI)	0.0 [0.0, 0.0]
9 Adverse events (local)	10		Risk Difference (M-H, Random, 95% CI)	Subtotals only
9.1 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMD	2	1652	Risk Difference (M-H, Random, 95% CI)	0.07 [0.04, 0.11]
9.2 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMV	2	516	Risk Difference (M-H, Random, 95% CI)	0.17 [0.01, 0.33]
9.3 Vitamin D vs. corticosteroid (very potent): calcipotriol vs. clobetasol propionate	2	194	Risk Difference (M-H, Random, 95% CI)	0.19 [0.10, 0.28]
9.4 Vitamin D + corticosteroid vs. corticosteroid: calcipotriol + BMD vs. BMD	3	2415	Risk Difference (M-H, Random, 95% CI)	-0.00 [-0.02, 0.01]
9.5 Vitamin D vs. vitamin D + corticosteroid: calcipotriol vs. calcipotriol + BMD	4	2801	Risk Difference (M-H, Random, 95% CI)	0.09 [0.06, 0.12]
9.6 Vitamin D vs. other treatments: calcipotriol vs. coal tar polytherapy	1	445	Risk Difference (M-H, Random, 95% CI)	0.24 [0.15, 0.33]
10 Adverse events (systemic)	6		Risk Difference (M-H, Random, 95% CI)	Subtotals only
10.1 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMD	2	1666	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.00, 0.00]
10.2 Vitamin D vs. corticosteroid (potent): calcipotriol vs. BMV	1	474	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]
10.3 Vitamin D vs. corticosteroid (very potent): calcipotriol vs. clobetasol propionate	1	151	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.03, 0.03]

10.4 Vitamin D + corticosteroid vs. corticosteroid:	2	2216	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.00, 0.00]
calcipotriol + BMD vs. BMD 10.5 Vitamin D vs. vitamin D + corticosteroid: calcipotriol vs.	3	1970	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.00, 0.00]
calcipotriol + BMD 10.6 Vitamin D vs. other treatments: calcipotriol vs. coal tar polytherapy	1	445	Risk Difference (M-H, Random, 95% CI)	0.0 [-0.01, 0.01]

Analysis I.I. Comparison I Vitamin D analogues versus placebo, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: I IAGI

itudy or subgroup Vitamir	n D analogue		Placebo		Std. Mean Difference	Weight	Std. Mean Difference
	N	Mean(SD)	N	Mean(SD)	IV,Random,95% Cl	V Velgite	IV,Random,95% CI
alcipotriol							
arker 1999 (P)	26	-3.19 (1.27)	28	-1.86 (0.89)	-	7.8 %	-1.20 [-1.79, -0.62]
Dubertret 1992	62	-2.66 (0.87)	62	-1.84 (0.75)	-	10.6 %	-1.00 [-1.38, -0.63]
leming 2010 (P)	74	2.3 (0.81)	28	2.64 (0.62)	-	9.7 %	-0.44 [-0.88, 0.00]
Guenther 2002 (P)	227	-3.3 (1.12)	206	-1.88 (1.14)	-	12.7 %	-1.25 [-1.46, -1.05]
larrington 1996a	290	-2.06 (0.8)	71	-1.37 (0.91)	-	12.0 %	-0.84 [-1.10, -0.57]
ang 1998	15	-3.87 (1.36)	15	-1.47 (0.99)	-	4.9 %	-1.96 [-2.86, -1.07]
aufmann 2002 (P)	480	2.2417 (0.9)	157	2.68 (0.89)	-	13.0 %	-0.49 [-0.67, -0.31]
ragballe 1988b	27	-2.4 (0.88)	27	-1.17 (0.62)	-	7.4 %	-1.59 [-2.21, -0.97]
Dranje 1997	43	-2.53 (1.05)	34	-1.91 (1.19)	-	9.4 %	-0.55 [-1.01, -0.09]
app 2003 (P)	308	-2.8 (.2)	107	-1.92 (1.07)	-	12.5 %	-0.76 [-0.98, -0.53]
ototal (95% CI)	1552		735		•	100.0 %	-0.93 [-1.17, -0.68]
erogeneity: Tau ² = 0.11; Chi ² = for overall effect: $Z = 7.37$ (P · alcipotriol plus occlusion		(P<0.00001); I	2 =82%				
ototal (95% CI) erogeneity: not applicable	0		0				Not estimable
for overall effect: not applicable alcitriol	e						

(Continued . . .)

Topical treatments for chronic plaque psoriasis (Review)

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(... Continued)

Study or subgroup	Vitamin D analogue		Placebo		Std. Mean Difference	Weight	Stc Mear Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% C
Langner 1992	29	-4.1 (1.05)	29	-3 (1.04)	-	16.2 %	-1.04 [-1.59, -0.49
Langner 1993	32	-3.97 (1.33)	32	-3.28 (1.14)	-	16.5 %	-0.55 [-1.05, -0.05
Langner 2001 (P)	15	-2.4 (1.18)	14	-2.14 (0.66)	-	14.9 %	-0.26 [-0.99, 0.47
Lebwohl 2007	202	-3.1 (1.4)	199	-2.42 (1.37)	-	17.8 %	-0.49 [-0.69, -0.29
Perez 1996	84	-3.2 (0.85)	84	-1.14 (0.38)	-	16.8 %	-3.11 [-3.57, -2.66
Powers 2005	199	-2.92 (1.27)	201	-2.01 (1.27)	-	17.8 %	-0.72 [-0.92, -0.5
Subtotal (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 4 Tacalcitol	65; Chi ² = 114.17, df =	5 (P<0.00001);	559 ² =96%		•	100.0 %	-1.03 [-1.71, -0.36
Langley 2011 (P)	163	2.28 (0.89)	64	2.75 (0.78)	•	50.0 %	-0.54 [-0.84, -0.25
Van de Kerkhof 1996a	a 103	-1.66 (0.88)	103	-0.75 (0.72)		50.0 %	-1.13 [-1.42, -0.83
Subtotal (95% CI) Heterogeneity: Tau ² = 0. Test for overall effect: Z = 5 Maxacalcitol Barker 1999 (P)	15; $Chi^2 = 7.57$, $df = 1$ (P = 0.01); I ² =	167 87% 28	-1.86 (0.89)	•	100.0 %	-0.84 [-1.41, -0.26
Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z = 6 Paricalcitol OD Durakovic 2004	cable	-2.45 (1.09)	28	-0.69 (0.94)	•	100.0 %	-1.43 [-1.91, -0.96 -1.66 [-2.66, -0.67
Heterogeneity: not applic Test for overall effect: Z =	cable	2.33 (0.84)	11 58	2.5 (0.68)	•	100.0 %	-1.66 [-2.66, -0.67
Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z =	cable = 3.27 (P = 0.0011) 63 63 cable = 1.21 (P = 0.23)	2.33 (0.84)		2.5 (0.68)	•		-0.22 [-0.58, 0.14
Heterogeneity: not applic Test for overall effect: Z = 7 Becocalcidiol OD Helfrich 2007 Subtotal (95% CI) Heterogeneity: not applic	cable = 3.27 (P = 0.0011) 63 63 cable = 1.21 (P = 0.23)	2.33 (0.84) 2.02 (0.74)	58	2.5 (0.68) 2.5 (0.68)	•	100.0 %	

Analysis I.2. Comparison I Vitamin D analogues versus placebo, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 2 TSS

Study or subgroup	Vitamin D analogue N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
l Calcipotriol							
Barker 1999 (P)	26	2.8 (2.3)	28	8.3 (3.2)		8.0 %	-1.93 [-2.59, -1.28]
Dubertret 1992	61	3.15 (1.94)	61	4.68 (1.94)	-	12.2 %	-0.78 [-1.15, -0.42]
Highton 1995	124	1.7 (1.2)	123	3.15 (1.2)	+	13.7 %	-1.20 [-1.48, -0.93]
Hinds n 2006 (P)	4	5.2 (3.3)	92	10.4 (4.4)	+	13.2 %	-1.35 [-1.66, -1.05]
Kang 1998	15	4.53 (3.33)	15	10.8 (3.33)		5.7 %	-1.83 [-2.70, -0.96]
Kragballe 1988b	27	4.63 (1.88)	27	6.73 (1.37)		8.8 %	-1.26 [-1.85, -0.67]
Levine 2010 (P)	48	3.52 (1.95)	48	4.08 (1.93)	-=-	11.6 %	-0.29 [-0.69, 0.12]
Pariser 1996	167	2.27 (1.33)	168	3.63 (1.33)	•	14.3 %	-1.02 [-1.25, -0.79]
Staberg 1989	9	1.3 (1.6)	9	4.6 (2.2)		4.1 %	-1.63 [-2.74, -0.53]
Staberg 1707							
Zonneveld 1998 (P) Subtotal (95% CI)	23 614	2.6 (1.73)	23 594	4.6 (1.73)	•	8.3 % 100.0 %	-1.14 [-1.76, -0.51] -1.15 [-1.41, -0.89]
Zonneveld 1998 (P) Subtotal (95% CI) Heterogeneity: Tau ² = 0.1 Test for overall effect: Z = 2 Calcipotriol plus occlusi	614 I I; Chi ² = 32.32, df = 9 = 8.66 (P < 0.00001) ion	(P = 0.00018);	594 ² =72%		•	100.0 %	-1.15 [-1.41, -0.89]
Zonneveld 1998 (P) Subtotal (95% CI) Heterogeneity: Tau ² = 0.1 Test for overall effect: Z = 2 Calcipotriol plus occlusi- Hinds n 2006 (P)	614 II; Chi ² = 32.32, df = 9 = 8.66 (P < 0.00001) ion 95		594 ² =72% 92	4.6 (1.73)	•	100.0 %	-1.15 [-1.41, -0.89] -0.15 [-0.44, 0.14]
Zonneveld 1998 (P) Subtotal (95% CI) Heterogeneity: Tau ² = 0.1 Test for overall effect: Z = 2 Calcipotriol plus occlusi	614 I I; Chi ² = 32.32, df = 9 = 8.66 (P < 0.00001) ion 95 95 able	(P = 0.00018);	594 ² =72%		•	100.0 %	
Zonneveld 1998 (P) Subtotal (95% CI) Heterogeneity: Tau ² = 0.1 Test for overall effect: Z = 2 Calcipotriol plus occlusi Hinds n 2006 (P) Subtotal (95% CI) Heterogeneity: not applica Test for overall effect: Z = 3 Calcitriol	614 II; Chi ² = 32.32, df = 9 = 8.66 (P < 0.00001) on 95 95 able = 1.03 (P = 0.30)	(P = 0.00018); 9.7 (4.8)	594 ² =72% 92 92	10.4 (4.4)	 • • •	100.0 % 100.0 % 100.0 %	-1.15 [-1.41, -0.89] -0.15 [-0.44, 0.14] -0.15 [-0.44, 0.14]
Zonneveld 1998 (P) Subtotal (95% CI) Heterogeneity: Tau ² = 0.1 Test for overall effect: Z = 2 Calcipotriol plus occlusi Hinds n 2006 (P) Subtotal (95% CI) Heterogeneity: not applica Test for overall effect: Z = 3 Calcitriol Lebwohl 2007	614 II; Chi ² = 32.32, df = 9 = 8.66 (P < 0.00001) ion 95 95 able = 1.03 (P = 0.30) 209	(P = 0.00018); 9.7 (4.8) 1.94 (0.89)	594 ² =72% 92 92 92	10.4 (4.4) 2.17 (0.83)	 • • • •	100.0 % 100.0 % 100.0 % 26.1 %	-1.15 [-1.41, -0.89] -0.15 [-0.44, 0.14] -0.15 [-0.44, 0.14] -0.27 [-0.46, -0.07]
Zonneveld 1998 (P) Subtotal (95% CI) Heterogeneity: Tau ² = 0.1 Test for overall effect: Z = 2 Calcipotriol plus occlusi Hinds n 2006 (P) Subtotal (95% CI) Heterogeneity: not applica Test for overall effect: Z = 3 Calcitriol Lebwohl 2007 Perez 1996	614 11; Chi ² = 32.32, df = 9 = 8.66 (P < 0.00001) ion 95 95 able = 1.03 (P = 0.30) 209 84	(P = 0.00018); 9.7 (4.8) 1.94 (0.89) 2.8 (1.5)	594 1 ² =72% 92 92 92 92 92 92 92 92 	10.4 (4.4) 2.17 (0.83) 7.1 (0.1)	 • • • •	100.0 % 100.0 % 100.0 % 26.1 % 24.9 %	-1.15 [-1.41, -0.89] -0.15 [-0.44, 0.14] -0.15 [-0.44, 0.14] -0.27 [-0.46, -0.07] -4.03 [-4.56, -3.50]

(Continued . . .)

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Vitamin D analogue N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	(Continued Std. Mean Difference IV,Random,95% Cl
Test for overall effect: Z =	2.07 (P = 0.038)						
4 Tacalcitol							
Scarpa 1997	134	3.44 (1.94)	134	4.34 (1.94)	-	47.8 %	-0.46 [-0.71, -0.22]
Seidenari 1997 (P)	11	2.92 (1.93)	П	4.52 (1.38)		9.5 %	-0.92 [-1.81, -0.03]
Van de Kerkhof 1996a	103	-4 (2.06)	103	-2.3 (2.06)	-	42.6 %	-0.82 [-1.11, -0.54]
Subtotal (95% CI)	248		248		•	100.0 %	-0.66 [-0.95, -0.36]
Heterogeneity: Tau ² = 0.02 Test for overall effect: Z = 5 Maxacalcitol Barker 1999 (P)		P = 0.14); I ² = 3.13 (3.24)	28	8.38 (3.2)	-	100.0 %	-1.61 [-2.10, -1.12]
Subtotal (95% CI)	75		28		•	100.0 %	-1.61 [-2.10, -1.12]
Heterogeneity: not applica Test for overall effect: Z = 6 Paricalcitol OD Durakovic 2004		2.7 (1.7)	11	7.6 (2.6)	-	100.0 %	-2.15 [-3.24, -1.06
Subtotal (95% CI) Heterogeneity: not applica Test for overall effect: Z = 7 Becocalcidiol OD			11		•	100.0 %	-2.15 [-3.24, -1.06]
Helfrich 2007	63	4.86 (2.23)	58	4.89 (1.62)		100.0 %	-0.02 [-0.37, 0.34]
Subtotal (95% CI) Heterogeneity: not applica Test for overall effect: Z = 8 Becocalcidiol twice daily	0.08 (P = 0.93)	. ,	58		•	100.0 %	-0.02 [-0.37, 0.34]
Helfrich 2007	61	4.03 (2.04)	58	4.89 (1.62)		100.0 %	-0.46 [-0.83, -0.10]
Subtotal (95% CI) Heterogeneity: not applica Test for overall effect: Z = Test for subgroup difference	2.49 (P = 0.013)	7 (P = 0.00), I ²	58 =89%		•	100.0 %	-0.46 [-0.83, -0.10]
				-4 Favours vitamin I	-2 0 2 D analogue Favours pl	4	

Analysis I.3. Comparison I Vitamin D analogues versus placebo, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 3 PASI

Study or subgroup Vi	tamin D analogue		Placebo		Std. Mean Difference	Weight	Std Mear Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% C
l Calcipotriol							
Dubertret 1992	65	6.3 (6.45)	65	9.16 (8.34)	-	9.2 %	-0.38 [-0.73, -0.03
Fleming 2010 (P)	74	4.3 (2.8)	28	5.5 (3.5)	-=-	6.6 %	-0.40 [-0.83, 0.04
Guenther 2002 (P)	227	4.2 (3.3)	207	7.7 (5.5)	-	17.2 %	-0.78 [-0.97, -0.58
Harrington 1996a	296	-4.3 (5.13)	70	-0.8 (5.4)	+	12.9 %	-0.67 [-0.94, -0.41
Kaufmann 2002 (P)	480	-0.46 (0.31)	157	-0.23 (0.34)	-	18.0 %	-0.72 [-0.91, -0.54
Mortensen 1993b	17	6.53 (2.49)	17	9.04 (4.71)		3.1 %	-0.65 [-1.34, 0.04
Oranje 1997	43	-0.52 (0.45)	34	-0.37 (0.4)	-=-	6.2 %	-0.35 [-0.80, 0.11
Papp 2003 (P)	308	-0.49 (0.32)	107	-0.29 (0.31)	-	15.3 %	-0.63 [-0.85, -0.40
Subtotal (95% CI)	1510		685		•	88.4 %	-0.65 [-0.75, -0.55
Heterogeneity: Tau ² = 0.00; C Test for overall effect: $Z = 12$ 2 Calcipotriol plus occlusion	.44 (P < 0.00001)						N
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol			0				Not estimabl
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not app 4 Tacalcitol			0				Not estimabl
Langley 2011 (P)	163	5.35 (4.44)	64	6.47 (3.56)	-	11.6 %	-0.27 [-0.56, 0.03
Subtotal (95% CI)	163		64		•	11.6 %	-0.27 [-0.56, 0.03
Heterogeneity: not applicable Test for overall effect: $Z = 1.7$ 5 Maxacalcitol							
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not app 6 Paricalcitol OD			0				Not estimabl
Subtotal (95% CI)	0		0				Not estimabl
				-4 Favours vitamin I		4 ebo	(Continued

Study or subgroup	Vitamin D analogue N	Placebo Mean(SD) N	Mean(SD)	S Me Differen IV,Random,9:	nce Weight	(Continued) Std. Mean Difference IV,Random,95% CI
Heterogeneity: not applica	able		· · /			
Test for overall effect: not						
7 Becocalcidiol OD	of the second seco					
Subtotal (95% CI)	0	0				Not estimable
Heterogeneity: not applica	able					
Test for overall effect: not	applicable					
8 Becocalcidiol twice daily	r					
Subtotal (95% CI)	0	0				Not estimable
Heterogeneity: not applica	able					
Test for overall effect: not	applicable					
Total (95% CI)	1673	749		•	100.0 %	-0.58 [-0.71, -0.45]
Heterogeneity: $Tau^2 = 0.0$	1; Chi ² = 13.87, df = 8	(P = 0.09); I ² =42%				
Test for overall effect: Z =	8.87 (P < 0.00001)					
Test for subgroup differen	ces: Chi ² = 5.94, df = 1	(P = 0.01), I ² =83%				
			_4	1 -2 0	2 4	
			Favours vitamin	D analogue F	avours placebo	

Analysis I.4. Comparison I Vitamin D analogues versus placebo, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 4 PAGI

Study or subgroup V	/itamin D analogue N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
l Calcipotriol					_		
Harrington 1996a	291	-2.27 (0.97)	71	-1.51 (1.07)	-	20.2 %	-0.77 [-1.03, -0.50]
Oranje 1997	43	-2.37 (1.02)	34	-1.91 (1.24)		10.7 %	-0.41 [-0.86, 0.05]
Subtotal (95% CI)	334		105		•	30.9 %	-0.64 [-0.97, -0.30]
Heterogeneity: $Tau^2 = 0.03$; Test for overall effect: $Z = 3$. 2 Calcipotriol plus occlusion	.67 (P = 0.00025)	$(P = 0.18); I^2 =$	-44%				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable Test for overall effect: not ap 3 Calcitriol							
Lebwohl 2007	202	-3.02 (1.37)	199	-2.32 (1.43)	-	25.4 %	-0.50 [-0.70, -0.30]
Powers 2005	199	-2.84 (1.33)	201	-1.95 (1.3)	-	25.1 %	-0.68 [-0.88, -0.47]
Subtotal (95% CI) Heterogeneity: Tau ² = 0.01;		(P = 0.22); I ² =	400		•	50.5 %	-0.59 [-0.76, -0.41]
Test for overall effect: $Z = 6$. 4 Tacalcitol		2.25 (0.02)		2.40.(0.00)			
Langley 2011 (P)	163	2.25 (0.93)	64	2.48 (0.99)		18.6 %	-0.24 [-0.53, 0.05]
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 1. 5 Maxacalcitol			64		•	18.6 %	-0.24 [-0.53, 0.05]
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not ap 6 Paricalcitol OD			0				Not estimable
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not ap			0				Not estimable
7 Becocalcidiol OD Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not ap			0				Not estimable
				-4 Favours vitamin		4 acebo	(Continued

Study or subgroup	Vitamin D analogue	Place	bo		Di	Std. Mean fference	Weight	(Continued) Std. Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	IV,Rand	lom,95% Cl		IV,Random,95% CI
8 Becocalcidiol twice daily	/							
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applic	able							
Test for overall effect: not	applicable							
Total (95% CI)	898	50	59		•		100.0 %	-0.54 [-0.72, -0.36]
Heterogeneity: $Tau^2 = 0.0$	02; Chi ² = 8.99, df = 4 (P = 0.06); I ² =56%						
Test for overall effect: Z =	= 6.02 (P < 0.00001)							
Test for subgroup differen	ces: $Chi^2 = 4.54$, $df = 2$	$(P = 0.10), I^2 = 56\%$						
							1	
				-4	-2	0 2	4	
				Favours vitamin [) analogue	Favours plac	cebo	

Analysis I.5. Comparison I Vitamin D analogues versus placebo, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D analogue		Placebo			Std. Mean rence	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI			IV,Random,95% CI
l Calcipotriol								
Barker 1999 (P)	26	-3.19 (1.27)	28	-1.86 (0.89)			4.7 %	-1.20 [-1.79, -0.62]
Dubertret 1992	62	-2.66 (0.87)	62	-1.84 (0.75)			6.5 %	-1.00 [-1.38, -0.63]
Fleming 2010 (P)	74	2.3 (0.81)	28	2.64 (0.62)	-		5.9 %	-0.44 [-0.88, 0.00]
Guenther 2002 (P)	227	-3.3 (1.12)	206	-1.88 (1.14)	•		7.8 %	-1.25 [-1.46, -1.05]
Harrington 1996a	290	-2.06 (0.8)	71	-1.37 (0.91)	+		7.4 %	-0.84 [-1.10, -0.57]
Highton 1995	124	1.7 (1.2)	123	3.15 (1.2)	+		7.3 %	-1.20 [-1.48, -0.93]
Hinds n 2006 (P)	4	5.2 (3.3)	92	10.4 (4.4)	-		7.1 %	-1.35 [-1.66, -1.05]
					-4 -2 0	2 4		
				Favours vitami	n D analogue	Favours place	bo	

(Continued . . .)

 ~0.49 [-0.67, -0.31 ~1.59 [-2.21, -0.97 ~0.29 [-0.69, 0.12 ~0.65 [-1.34, 0.04 	2.9 % 8.0 %	[N	Mean(SD)	N	
 ~ -1.59 [-2.21, -0.97 ~ -0.29 [-0.69, 0.12 ~ -0.65 [-1.34, 0.04 	8.0 %		-1.47 (0.99)	15	-3.87 (1.36)	15	Kang 1998
 ~ -0.29 [-0.69, 0.12 ~ -0.65 [-1.34, 0.04 		-	2.68 (0.89)	157	2.2417 (0.9)	480	Kaufmann 2002 (P)
% -0.65 [-1.34, 0.04	4.5 %		-1.17 (0.62)	27	-2.4 (0.88)	27	Kragballe 1988b
	6.2 %		4.08 (1.93)	48	3.52 (1.95)	48	Levine 2010 (P)
% -0.55 [-1.01, -0.09	4.0 %		9.04 (4.71)	17	6.53 (2.49)	17	Mortensen 1993b
	5.7 %		-1.91 (1.19)	34	-2.53 (1.05)	43	Oranje 1997
% -0.76 [-0.98, -0.53	7.7 %	+	-1.92 (1.07)	107	-2.81 (1.21)	308	Papp 2003 (P)
% -1.02 [-1.25, -0.79	7.7 %	-	3.63 (1.33)	168	2.27 (1.33)	167	Pariser 1996
% -1.63 [-2.74, -0.53	2.2 %		4.6 (2.2)	9	1.3 (1.6)	9	Staberg 1989
% -1.14 [-1.76, -0.51	4.4 %		4.6 (1.73)	23	2.6 (1.73)	23	Zonneveld 1998 (P)
% - 0.96 [-1.15, -0.77	100.0 %	•		1215 ² =79%	6 (P<0.00001);	= 9.98 (P < 0.00001)	Subtotal (95% CI) Heterogeneity: Tau ² = 0.1 Test for overall effect: Z = 2 Calcipotriol plus occlusio
% -0.15 [-0.44, 0.14	100.0 %	-	10.4 (4.4)	92	9.7 (4.8)	95	Hinds n 2006 (P)
% -0.15 [-0.44, 0.14	100.0 %	•		92			Subtotal (95% CI) Heterogeneity: not applica Test for overall effect: Z = 3 Calcitriol
% -1.04 [-1.59, -0.49	14.2 %		-3 (1.04)	29	-4.1 (1.05)	29	Langner 1992
% -0.55 [-1.05, -0.05	14.5 %		-3.28 (1.14)	32	-3.97 (1.33)	32	Langner 1993
% -0.26 [-0.99, 0.47	3. %		-2.14 (0.66)	14	-2.4 (1.18)	15	Langner 2001 (P)
% -0.49 [-0.69, -0.29	15.7 %	-	-2.42 (1.37)	199	-3.1 (1.4)	202	Lebwohl 2007
% -3.11 [-3.57, -2.66	14.7 %		-1.14 (0.38)	84	-3.2 (0.85)	84	Perez 1996
% -0.72 [-0.92, -0.51	15.7 %	-	-2.01 (1.27)	201	-2.92 (1.27)	199	Powers 2005
% -0.06 [-0.94, 0.81	12.1 %		6.1 (1.49)	10	6 (1.49)	10	Van de Kerkhof 1989
% -0.92 [-1.54, -0.29	100.0 %	•		569 ² =95%	6 (P<0.00001);		Subtotal (95% CI) Heterogeneity: Tau ² = 0.6 Test for overall effect: Z = 4 Tacalcitol
% -0.54 [-0.84, -0.25	29.0 %	-	2.75 (0.78)	64	2.28 (0.89)	163	Langley 2011 (P)
% -0.46 [-0.71, -0.22	30.8 %	-	4.34 (1.94)	134	3.44 (1.94)	134	Scarpa 1997
% -0.92 [-1.81, -0.03	11.3 %		4.52 (1.38)	11	2.92 (1.93)	11	Seidenari 1997 (P)
% -1.13 [-1.42, -0.83	28.9 %	-	-0.75 (0.72)	103	-1.66 (0.88)	103	Van de Kerkhof 1996a

Study or subgroup	Vitamin D analogue N	F Mean(SD)	Placebo N	Mean(SD)		Std. Mean erence m,95% Cl	Weight	(Continued Std. Mean Difference IV,Random,95% CI
Subtotal (95% CI)	411		312		•		100.0 %	-0.73 [-1.09, -0.37]
Heterogeneity: $Tau^2 = 0.0$	9; Chi ² = 13.06, df = 3	$(P = 0.005); I^2 =$	77%					
Test for overall effect: Z =	3.98 (P = 0.000069)							
5 Maxacalcitol					_			
Barker 1999 (P)	75	-3.52 (1.23)	28	-1.86 (0.89)			100.0 %	-1.43 [-1.91, -0.96]
Subtotal (95% CI)	75		28		•		100.0 %	-1.43 [-1.91, -0.96]
Heterogeneity: not applica	able							
Test for overall effect: Z =	5.88 (P < 0.00001)							
6 Paricalcitol OD								
Durakovic 2004	11	-2.45 (1.09)	11	-0.69 (0.94)			100.0 %	-1.66 [-2.66, -0.67]
Subtotal (95% CI)	11		11		-		100.0 %	-1.66 [-2.66, -0.67]
Heterogeneity: not applica	able							
Test for overall effect: Z =	3.27 (P = 0.0011)							
7 Becocalcidiol OD								
Helfrich 2007	63	2.33 (0.84)	58	2.5 (0.68)			100.0 %	-0.22 [-0.58, 0.14]
Subtotal (95% CI)	63		58		•		100.0 %	-0.22 [-0.58, 0.14]
Heterogeneity: not applica	able							
Test for overall effect: $Z =$	· /							
8 Becocalcidiol twice daily					_			
Helfrich 2007	61	2.02 (0.74)	58	2.5 (0.68)			100.0 %	-0.67 [-1.04, -0.30]
Subtotal (95% CI)	61		58		•		100.0 %	-0.67 [-1.04, -0.30]
Heterogeneity: not applica	able							
Test for overall effect: Z =	· · · · ·							
Test for subgroup difference	ces: $Chi^2 = 41.16$, df =	7 (P = 0.00), $I^2 =$	83%					
							1	
				-4	-2 0	2	4	
				Favours vitamin D) analogue	Favours pla	cebo	

Analysis I.6. Comparison I Vitamin D analogues versus placebo, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 6 Total withdrawals

H,Random,95% H,Random,95% 1/N n/N 4/30 2/30 4/66 4/66 4/66 4/66 24/214 14/109 19/223 15/113 6/79 12/40		
4/66 4/66 4.6 % 0.0 [-0.08, 0.08] 24/214 14/109 4.9 % -0.02 [-0.09, 0.06] 19/223 15/113 5.1 % -0.05 [-0.12, 0.02]		
4/66 4/66 4.6 % 0.0 [-0.08, 0.08] 24/214 14/109 4.9 % -0.02 [-0.09, 0.06] 19/223 15/113 5.1 % -0.05 [-0.12, 0.02]		l Calcipotriol
24/214 14/109 4.9 % -0.02 [-0.09, 0.06] 19/223 15/113 5.1 % -0.05 [-0.12, 0.02]	4	Barker 1999 (P)
19/223 15/113 5 .1% -0.05 [-0.12, 0.02]	2	Dubertret 1992
	24/	Feldman 2010 (1)
6/79 12/40 2.0 % -0.22 [-0.38, -0.07]	19/	Feldman 2010 (2)
	é	Fleming 2010 (P)
23/231 34/208 5.7 % -0.06 [-0.13, 0.00]	23/	Guenther 2002 (P)
30/326 17/87 + 4.2 % -0.10 [-0.19, -0.01]	30/	Harrington 1996a
0/15 0/15 2.9 % 0.0 [-0.12, 0.12]	(Kang 1998
39/480 25/157 • 5.7 % -0.08 [-0.14, -0.02]	39/	Kaufmann 2002 (P)
2/48 0/48 5.4 % 0.04 [-0.03, 0.11]	2	Levine 2010 (P)
0/17 0/17 - 3.3 % 0.0 [-0.11, 0.11]	(Mortensen 1993b
6/43 3/34 2.3 % 0.05 [-0.09, 0.19]	é	Oranje 1997
27/308 12/108 5.4 % -0.02 [-0.09, 0.04]	27/	Рарр 2003 (Р)
0/10 0/10 1.6 % 0.0 [-0.17, 0.17]	(Staberg 1989
2090 1042 55.0 % -0.03 [-0.06, 0.00]	20	Subtotal (95% CI)
$3.15, df = 13 (P = 0.04); l^2 = 44\%$ 0.066) 0 0 Not estimable	= 13	Total events: 184 (Vitamin D analogue), 13 Heterogeneity: Tau ² = 0.00; Chi ² = 23.15, Test for overall effect: Z = 1.84 (P = 0.066) 2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D analogue), 0 (Pla Heterogeneity: not applicable
		Test for overall effect: not applicable
		3 Calcitriol
1/32 1/32 4.3 % 0.0 [-0.09, 0.09]		Langner 1992
0/29 0/29 5.6 % 0.0 [-0.06, 0.06]	(Langner 1993
1/15 3/14 -0.15 [-0.40, 0.10]		Langner 2001 (P)

(... Continued)

Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	IM- H,Random,95% Cl		M- H,Random, C
Perez 1996	0/84	0/84	-	8.3 %	0.0 [-0.02, 0.02]
Van de Kerkhof 1989	0/10	0/10		1.6 %	0.0 [-0.17, 0.17]
Subtotal (95% CI)	170	169	•	20.8 %	0.00 [-0.02, 0.02]
Total events: 2 (Vitamin D ana Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.1 4 Tacalcitol	$hi^2 = 2.53$, df = 4 (P = 0.64);	l ² =0.0%			
Langley 2011 (P)	21/184	27/91		3.5 %	-0.18 [-0.29, -0.08]
Scarpa 1997	23/157	23/157	+	4.7 %	0.0 [-0.08, 0.08]
Seidenari 1997 (P)	1/12	1/12		1.1 %	0.0 [-0.22, 0.22]
Van de Kerkhof 1996a	19/122	19/122	+	4.1 %	0.0 [-0.09, 0.09]
Subtotal (95% CI)	475	382	•	13.3 %	-0.05 [-0.14, 0.05]
Heterogeneity: Tau ² = 0.01; C Test for overall effect: Z = 1.0 5 Maxacalcitol Barker 1999 (P)	,	2/30		2.9 %	0.11 [-0.01, 0.23
Subtotal (95% CI)	90	30	•	2.9 %	0.11 [-0.01, 0.23]
Total events: 16 (Vitamin D ar Heterogeneity: not applicable Test for overall effect: Z = 1.8 6 Paricalcitol OD Durakovic 2004		0/11	_	1.9 %	0.0 [-0.16, 0.16]
Subtotal (95% CI)	11	11	+	1.9 %	0.0 [-0.16, 0.16]
Fotal events: 0 (Vitamin D ana Heterogeneity: not applicable Fest for overall effect: Z = 0.0 7 Becocalcidiol OD Helfrich 2007	- / . /	53/60	_	3.1 %	-0.01 [-0.12, 0.11]
Subtotal (95% CI)	64	60	+	3.1 %	-0.01 [-0.12, 0.11]
Total events: 56 (Vitamin D ar Heterogeneity: not applicable		53/60		3.0 %	-0.01 [-0.13, 0.10
Becocalcidiol twice daily	53/61			5.0 /0	0.01 [-0.10, 0.10]
Test for overall effect: Z = 0.1 Becocalcidiol twice daily Helfrich 2007 Subtotal (95% CI)	53/61 61	60	•	3.0 %	-0.01 [-0.13, 0.10]

(Continued . . .)

Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M-	Weight	(Continued) Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
Total (95% CI)	2961	1754	•	100.0 %	-0.02 [-0.05, 0.00]
Total events: 375 (Vitamin E) analogue), 320 (Placebo)				
Heterogeneity: $Tau^2 = 0.00$;	Chi ² = 53.54, df = 26 (P = 0.0	001); I ² =51%			
Test for overall effect: $Z = I$.71 (P = 0.087)				
Test for subgroup difference	es: $Chi^2 = 6.83$, $df = 6$ (P = 0.34	4), I ² = I 2%			
			-1 -0.5 0 0.5	I	
		Favours vitar	nin D analogue Favours plac	cebo	

Analysis I.7. Comparison I Vitamin D analogues versus placebo, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
l Calcipotriol					
Barker 1999 (P)	0/30	1/30	-	1.7 %	-0.03 [-0.12, 0.05]
Dubertret 1992	2/66	1/66		4.1 %	0.02 [-0.04, 0.07]
Feldman 2010 (1)	7/214	1/109	•	7.7 %	0.02 [-0.01, 0.05]
Feldman 2010 (2)	2/223	3/113	•	7.2 %	-0.02 [-0.05, 0.01]
Guenther 2002 (P)	6/227	21/208	+	4.7 %	-0.07 [-0.12, -0.03]
Harrington 1996a	8/326	8/174	•	6.5 %	-0.02 [-0.06, 0.01]
Highton 1995	6/139	8/138	+	4.0 %	-0.01 [-0.07, 0.04]
Kang 1998	0/15	0/15	-	1.0 %	0.0 [-0.12, 0.12]
Kaufmann 2002 (P)	15/480	12/157	+	5.0 %	-0.05 [-0.09, 0.00]
			-I -0.5 0 0.5 I		
		Favours vi	tamin D analogue Favours placebo		

(Continued . . .)

			Risk		(Continued Risk
Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M-	Weight	Nisk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
Levine 2010 (P)	0/48	0/48		5.7 %	0.0 [-0.04, 0.04]
Mortensen 1993b	0/17	0/17	-	1.2 %	0.0 [-0.11, 0.11]
Staberg 1989	0/10	0/10		0.5 %	0.0 [-0.17, 0.17]
Subtotal (95% CI)	1795	1085		49.2 %	-0.02 [-0.04, 0.00]
Fotal events: 46 (Vitamin D ;		1085		49.2 %	-0.02 [-0.04, 0.00]
	$Chi^2 = 20.47, df = 11 (P = 0.0)$	04); l ² =46%			
Test for overall effect: $Z = 1$.		,. ,.			
2 Calcipotriol plus occlusion					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D ar	nalogue), 0 (Placebo)				
Heterogeneity: not applicabl	e				
Test for overall effect: not ap	plicable				
3 Calcitriol					
Langner 1992	0/29	0/29	-	2.9 %	0.0 [-0.06, 0.06]
Langner 1993	1/32	1/32	+	1.8 %	0.0 [-0.09, 0.09]
Langner 2001 (P)	1/15	3/14	- _	0.2 %	-0.15 [-0.40, 0.10]
Perez 1996	0/84	0/84	•	9.4 %	0.0 [-0.02, 0.02]
Van de Kerkhof 1989	0/10	0/10		0.5 %	0.0 [-0.17, 0.17
Subtotal (95% CI)	170	169	•	14.8 %	0.00 [-0.02, 0.02]
Total events: 2 (Vitamin D ar	nalogue), 4 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$; C	$Chi^2 = 2.53$, df = 4 (P = 0.64);	$ ^2 = 0.0\%$			
Test for overall effect: $Z = 0$.	.10 (P = 0.92)				
4 Tacalcitol					
Langley 2011 (P)	4/184	4/91	-	4.6 %	-0.02 [-0.07, 0.02]
Scarpa 1997	1/157	0/157	a	11.0 %	0.01 [-0.01, 0.02]
Seidenari 1997 (P)	0/12	0/12		0.7 %	0.0 [-0.15, 0.15]
Van de Kerkhof 1996a	1/122	0/122	•	9.6 %	0.01 [-0.01, 0.03]
Subtotal (95% CI)	475	382	•	25.8 %	0.00 [-0.01, 0.02]
Total events: 6 (Vitamin D ar	nalogue), 4 (Placebo)				
. ,	$Chi^2 = 2.30, df = 3 (P = 0.51);$	$ ^2 = 0.0\%$			
Test for overall effect: $Z = 0$.	.70 (P = 0.48)				
5 Maxacalcitol					
Barker 1999 (P)	2/90	1/30	+	2.5 %	-0.01 [-0.08, 0.06]
Subtotal (95% CI)	90	30	•	2.5 %	-0.01 [-0.08, 0.06]
Total events: 2 (Vitamin D ar	nalogue), I (Placebo)				
Heterogeneity: not applicabl					
Test for overall effect: $Z = 0$.	.31 (P = 0.76)				
6 Paricalcitol OD					

- | -0.5 0 0.5 1

Favours vitamin D analogue Favours placebo

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Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M- H,Random,95%	Weight	Risk Difference M- H,Random,95:
	n/N	n/N	CI		H,Rahdom,75. Cl
Durakovic 2004	0/11	0/11	+	0.6 %	0.0 [-0.16, 0.16]
Subtotal (95% CI)	11	11	+	0.6 %	0.0 [-0.16, 0.16]
Total events: 0 (Vitamin D analo	ogue), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 0.0 (P = 1.0)				
7 Becocalcidiol OD					
Helfrich 2007	2/64	0/60	-+-	4.0 %	0.03 [-0.02, 0.08]
Subtotal (95% CI)	64	60	•	4.0 %	0.03 [-0.02, 0.08]
Total events: 2 (Vitamin D analo	ogue), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 1.18	(P = 0.24)				
8 Becocalcidiol twice daily					
Helfrich 2007	3/61	0/60	+	3.1 %	0.05 [-0.01, 0.11]
Subtotal (95% CI)	61	60	•	3.1 %	0.05 [-0.01, 0.11]
Total events: 3 (Vitamin D analo	ogue), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 1.56	(P = 0.12)				
Total (95% CI)	2666	1797		100.0 %	0.00 [-0.02, 0.01]
Total events: 61 (Vitamin D ana	llogue), 64 (Placebo)				
Heterogeneity: $Tau^2 = 0.00$; Ch	$i^2 = 37.74$, df = 24 (P = 0.0	04); I ² =36%			
Test for overall effect: $Z = 0.73$	(P = 0.47)				
Test for subgroup differences: C	$Chi^2 = 6.56, df = 6 (P = 0.3)$	6), l ² =9%			

Favours vitamin D analogue Favours placebo

Analysis I.8. Comparison I Vitamin D analogues versus placebo, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
I Calcipotriol					
Barker 1999 (P)	0/30	1/30	-	4.9 %	-0.03 [-0.12, 0.05]
Feldman 2010 (1)	4/214	3/109	-	8.8 %	-0.01 [-0.04, 0.03]
Feldman 2010 (2)	3/223	1/113	+	9.8 %	0.00 [-0.02, 0.03]
Guenther 2002 (P)	2/227	19/208	-	8.4 %	-0.08 [-0.12, -0.04]
Harrington 1996a	9/326	22/174	-	7.4 %	-0.10 [-0.15, -0.05]
Levine 2010 (P)	0/48	0/48	+	8.5 %	0.0 [-0.04, 0.04]
Staberg 1989	0/10	0/10		1.9 %	0.0 [-0.17, 0.17]
Subtotal (95% CI)	1078	692	•	49.6 %	-0.03 [-0.08, 0.01]
Test for overall effect: $Z = 1.4$	J(1 - 0.15)				
Test for overall effect: Z = 1.4 2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable	0 alogue), 0 (Placebo)	0			Not estimable
2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app	O alogue), 0 (Placebo)	0			Not estimable
2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol	O alogue), 0 (Placebo)	0		5.2 %	
2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app	O alogue), 0 (Placebo) licable		-	5.2 %	Not estimable
2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol Langner 1993	0 alogue), 0 (Placebo) licable 0/32	1/32			-0.03 [-0.11, 0.05]
2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol Langner 1993 Langner 2001 (P)	0 alogue), 0 (Placebo) licable 0/32 0/15	1/32 3/14		1.2 %	-0.03 [-0.11, 0.05] -0.21 [-0.44, 0.02]
2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol Langner 1993 Langner 2001 (P) Perez 1996 Van de Kerkhof 1989 Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: Tau ² = 0.00; C	0 alogue), 0 (Placebo) llicable 0/32 0/15 0/84 0/10 141 alogue), 4 (Placebo) Chi ² = 8.79, df = 3 (P = 0.03	1/32 3/14 0/84 0/10 140		1.2 % 9.8 %	-0.03 [-0.11, 0.05] -0.21 [-0.44, 0.02] 0.0 [-0.02, 0.02]
2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol Langner 1993 Langner 2001 (P) Perez 1996	0 alogue), 0 (Placebo) llicable 0/32 0/15 0/84 0/10 141 alogue), 4 (Placebo) Chi ² = 8.79, df = 3 (P = 0.03	1/32 3/14 0/84 0/10 140		1.2 % 9.8 % 1.9 %	-0.03 [-0.11, 0.05] -0.21 [-0.44, 0.02] 0.0 [-0.02, 0.02] 0.0 [-0.17, 0.17]

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					(Continue
Study or subgroup	Vitamin D analogue	Placebo	Risk Difference	Weight	Risk Difference
/8			M- H,Random,95%		M. H,Random,
	n/N	n/N	H,Random,25% Cl		H,Random, C
Total events: 0 (Vitamin D ar	nalogue), 0 (Placebo)				
Heterogeneity: not applicable	e				
Test for overall effect: $Z = 0$.	0 (P = 1.0)				
5 Maxacalcitol					
Barker 1999 (P)	0/90	1/30	-	5.5 %	-0.03 [-0.11, 0.04
Subtotal (95% CI)	90	30	•	5.5 %	-0.03 [-0.11, 0.04]
Total events: 0 (Vitamin D ar	nalogue), I (Placebo)				
Heterogeneity: not applicable	e				
Test for overall effect: $Z = 0$.	85 (P = 0.40)				
6 Paricalcitol OD					
Durakovic 2004	0/11	0/11	+	2.1 %	0.0 [-0.16, 0.16
Subtotal (95% CI)	11	11	+	2.1 %	0.0 [-0.16, 0.16]
Total events: 0 (Vitamin D ar	nalogue), 0 (Placebo)				
Heterogeneity: not applicable	e				
Test for overall effect: $Z = 0$.	0 (P = 1.0)				
7 Becocalcidiol OD					
Helfrich 2007	1/64	2/60	-	7.2 %	-0.02 [-0.07, 0.04
Subtotal (95% CI)	64	60	•	7.2 %	-0.02 [-0.07, 0.04]
Total events: I (Vitamin D ar	nalogue), 2 (Placebo)				
Heterogeneity: not applicable	e				
Test for overall effect: $Z = 0$.	.64 (P = 0.53)				
8 Becocalcidiol twice daily					
Helfrich 2007	0/61	2/60	-	7.2 %	-0.03 [-0.09, 0.02
Subtotal (95% CI)	61	60	•	7.2 %	-0.03 [-0.09, 0.02
Total events: 0 (Vitamin D ar	nalogue), 2 (Placebo)				
Heterogeneity: not applicable	JJJJJJJJJJJJJ				
Test for overall effect: $Z = 1$.					
Total (95% CI)	1602	1150	•	100.0 %	-0.03 [-0.05, 0.00
Total events: 19 (Vitamin D a	analogue), 55 (Placebo)				
Heterogeneity: $Tau^2 = 0.00;$	Chi ² = 81.93, df = 15 (P<0.0	0001); I ² =82%			
Test for overall effect: $Z = 1$.	93 (P = 0.054)				
Test fen sub men in differen se	s: Chi ² = 4.30, df = 6 (P = 0.6	4) $ ^2 = 0.0\%$			

- I -0.5 0 0.5 I Favours vitamin D analogue Favours placebo

Analysis I.9. Comparison I Vitamin D analogues versus placebo, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9! Cl
l Calcipotriol					
Dubertret 1992	14/66	16/66		1.0 %	-0.03 [-0.17, 0.11]
Fleming 2010 (P)	8/79	10/40	<u>← ; </u>	0.9 %	-0.15 [-0.30, 0.00]
Guenther 2002 (P)	45/227	26/208	·	4.4 %	0.07 [0.00, 0.14]
Harrington 1996a	81/326	20/87		2.1 %	0.02 [-0.08, 0.12]
Highton 1995	28/139	21/138		2.6 %	0.05 [-0.04, 0.14]
Kang 1998	2/15	0/15		0.5 %	0.13 [-0.06, 0.33]
Kaufmann 2002 (P)	54/480	21/157		5.7 %	-0.02 [-0.08, 0.04]
Levine 2010 (P)	9/48	11/48		0.8 %	-0.04 [-0.20, 0.12]
Oranje 1997	7/43	8/34	· · · · · · · · · · · · · · · · · · ·	0.6 %	-0.07 [-0.25, 0.11]
Papp 2003 (P)	53/308	17/108		3.2 %	0.01 [-0.07, 0.10]
Staberg 1989	1/10	1/10	·	0.3 %	0.0 [-0.26, 0.26]
Subtotal (95% CI)	1741	911	-	22.3 %	0.01 [-0.03, 0.05]
Heterogeneity: Tau ² = 0.00; C Test for overall effect: Z = 0.4 2 Calcipotriol plus occlusion Subtotal (95% CI)	8 (P = 0.63) 0	.26); ² = 9% 0			Not estimable
Total events: 0 (Vitamin D ana	- , , , ,				
Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol					
Test for overall effect: not app		0/29		5.0 %	0.0 [-0.06, 0.06]
Test for overall effect: not app 3 Calcitriol	licable	0/29		5.0 %	0.0 [-0.06, 0.06] -0.01 [-0.06, 0.03]
Test for overall effect: not app 3 Calcitriol Langner 1992	licable 0/29		 		
Test for overall effect: not app 3 Calcitriol Langner 1992 Lebwohl 2007	olicable 0/29 11/209	14/209		10.1 %	-0.01 [-0.06, 0.03]

(Continued ...)

Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M-	Weight	Risk Difference M
	n/N	n/N	H,Random,95% Cl		H,Random, C
Heterogeneity: $Tau^2 = 0.0$; C	$hi^2 = 0.18, df = 3 (P = 0.98);$	$ ^2 = 0.0\%$			
Test for overall effect: $Z = 0.5$	51 (P = 0.61)				
4 Tacalcitol					
Langley 2011 (P)	32/184	4/9		2.5 %	0.02 [-0.07, 0.11
Scarpa 1997	1/134	2/157	-	40.1 %	-0.01 [-0.03, 0.02
Subtotal (95% CI)	318	248	+	42.5 %	0.00 [-0.03, 0.03]
Total events: 33 (Vitamin D a	nalogue), 16 (Placebo)				
Heterogeneity: Tau ² = 0.00; ($Chi^2 = 1.08, df = 1 (P = 0.30)$); I ² =7%			
Test for overall effect: Z = 0.2	21 (P = 0.83)				
5 Maxacalcitol					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D an	alogue), 0 (Placebo)				
Heterogeneity: not applicable	2				
Test for overall effect: not app	olicable				
6 Paricalcitol OD					
Durakovic 2004	0/11	0/11		0.8 %	0.0 [-0.16, 0.16
Subtotal (95% CI)	11	11		0.8 %	0.0 [-0.16, 0.16]
Total events: 0 (Vitamin D an	alogue), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 0.0	0 (P = 1.0)				
7 Becocalcidiol OD					
Helfrich 2007	3/64	2/60		4.4 %	0.01 [-0.06, 0.08
Subtotal (95% CI)	64	60	-	4.4 %	0.01 [-0.06, 0.08]
Total events: 3 (Vitamin D an	alogue), 2 (Placebo)				
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.3$	39 (P = 0.70)				
8 Becocalcidiol twice daily					
Helfrich 2007	8/61	2/60		2.3 %	0.10 [0.00, 0.19
Subtotal (95% CI)	61	60		2.3 %	0.10 [0.00, 0.19]
Total events: 8 (Vitamin D an	alogue), 2 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 1.9$	99 (P = 0.046)				
Total (95% CI)	2653	1749	+	100.0 %	0.00 [-0.01, 0.02]
Total events: 367 (Vitamin D					
Heterogeneity: $Tau^2 = 0.0$; C	$hi^2 = 18.36$, $df = 19$ (P = 0.5	0); l ² =0.0%			
Test for overall effect: $Z = 0.2$	21 (P = 0.83)				
Test for subgroup differences	$: Chi^2 = 4.6I, df = 5 (P = 0.4)$	6), l ² =0.0%			

Favours vitamin D analogue Favours placebo

(... Continued)

Analysis I.10. Comparison I Vitamin D analogues versus placebo, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: I Vitamin D analogues versus placebo

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D analogue	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,959 Cl
l Calcipotriol					
Barker 1999 (P)	0/30	0/30	-	0.8 %	0.0 [-0.06, 0.06]
Dubertret 1992	0/66	0/66	-	3.7 %	0.0 [-0.03, 0.03]
Harrington 1996a	0/326	0/87	-	11.9 %	0.0 [-0.02, 0.02]
Kang 1998	0/15	0/15	-	0.2 %	0.0 [-0.12, 0.12]
Mortensen 1993b	0/17	0/17	-	0.3 %	0.0 [-0.11, 0.11]
Oranje 1997	0/43	0/34	-	1.3 %	0.0 [-0.05, 0.05]
Papp 2003 (P)	0/308	0/108	-	17.5 %	0.0 [-0.01, 0.01]
Staberg 1989	0/10	0/10	—	0.1 %	0.0 [-0.17, 0.17]
Subtotal (95% CI)	815	367		35.8 %	0.0 [-0.01, 0.01]
Test for overall effect: $Z = 0.0$ 2 Calcipotriol plus occlusion	$hi^2 = 0.0, df = 7 (P = 1.00); I^2$ 0 (P = 1.0)				Not estimable
Heterogeneity: Tau ² = 0.0; C Test for overall effect: Z = 0.0 2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D an Heterogeneity: not applicable Test for overall effect: not appl 3 Calcitriol	hi ² = 0.0, df = 7 (P = 1.00); l ² 0 (P = 1.0) 0 halogue), 0 (Placebo)	0		0.0 %	Not estimable
Heterogeneity: Tau ² = 0.0; C Test for overall effect: Z = 0.1 2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D an Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol Langner 1992	hi ² = 0.0, df = 7 (P = 1.00); l ² 0 (P = 1.0) alogue), 0 (Placebo) plicable 0/29	0 0/29	-	0.8 %	0.0 [-0.06, 0.06]
Heterogeneity: Tau ² = 0.0; C Test for overall effect: Z = 0.0 2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D an Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol Langner 1992 Perez 1996	hi ² = 0.0, df = 7 (P = 1.00); l ² 0 (P = 1.0) alogue), 0 (Placebo) plicable 0/29 0/84	0 0/29 0/84		6.0 %	0.0 [-0.06, 0.06] 0.0 [-0.02, 0.02]
Heterogeneity: Tau ² = 0.0; C Test for overall effect: Z = 0.1 2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D an Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol Langner 1992	hi ² = 0.0, df = 7 (P = 1.00); l ² 0 (P = 1.0) alogue), 0 (Placebo) plicable 0/29	0 0/29	-		0.0 [-0.06, 0.06]
Heterogeneity: $Tau^2 = 0.0$; C Test for overall effect: $Z = 0.0$ 2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D an Heterogeneity: not applicable Test for overall effect: not app 3 Calcitriol Langner 1992 Perez 1996 Powers 2005 Subtotal (95% CI) Total events: 0 (Vitamin D an Heterogeneity: Tau ² = 0.0; C Test for overall effect: $Z = 0.0$ 4 Tacalcitol	hi ² = 0.0, df = 7 (P = 1.00); l ² 0 (P = 1.0) 0 halogue), 0 (Placebo) plicable 0/29 0/84 0/210 323 halogue), 0 (Placebo) hi ² = 0.0, df = 2 (P = 1.00); l ² 0 (P = 1.0)	0 0/29 0/84 0/211 324 =0.0%	-	6.0 % 36.9 % 43.6 %	0.0 [-0.06, 0.06] 0.0 [-0.02, 0.02] 0.0 [-0.01, 0.01] 0.0 [-0.01, 0.01]
Heterogeneity: Tau ² = 0.0; C Test for overall effect: Z = 0.0 2 Calcipotriol plus occlusion Subtotal (95% CI) Total events: 0 (Vitamin D an Heterogeneity: not applicable Test for overall effect: not ap 3 Calcitriol Langner 1992 Perez 1996 Powers 2005 Subtotal (95% CI) Total events: 0 (Vitamin D an	hi ² = 0.0, df = 7 (P = 1.00); l ² 0 (P = 1.0) 0 alogue), 0 (Placebo) 0 0/29 0/84 0/210 323 alogue), 0 (Placebo) hi ² = 0.0, df = 2 (P = 1.00); l ²	0 0/29 0/84 0/211 324	-	6.0 % 36.9 %	0.0 [-0.06, 0.06] 0.0 [-0.02, 0.02] 0.0 [-0.01, 0.01]

-1 -0.5 0 0.5 I Favours vitamin D analogue Favours placebo

(Continued ...)

Risk Risk Difference M-Difference Study or subgroup Vitamin D analogue Placebo Weight M-H,Random,95% H,Random,95% n/N n/N ĆΓ ĆΙ Total events: 0 (Vitamin D analogue), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0) 5 Maxacalcitol Barker 1999 (P) 0.0 [-0.05, 0.05] 0/90 0/30 1.4 % Subtotal (95% CI) 90 30 1.4 % 0.0 [-0.05, 0.05] Total events: 0 (Vitamin D analogue), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0) 6 Paricalcitol OD Durakovic 2004 0/11 0/11 0.1 % 0.0 [-0.16, 0.16] Subtotal (95% CI) 11 11 0.1 % 0.0 [-0.16, 0.16] Total events: 0 (Vitamin D analogue), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0) 7 Becocalcidiol OD Helfrich 2007 0/64 0/60 3.3 % 0.0 [-0.03, 0.03] Subtotal (95% CI) 64 60 3.3 % 0.0 [-0.03, 0.03] Total events: 0 (Vitamin D analogue), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0) 8 Becocalcidiol twice daily Helfrich 2007 0/64 0/60 3.3 % 0.0 [-0.03, 0.03] Subtotal (95% CI) 64 3.3 % 0.0 [-0.03, 0.03] 60 Total events: 0 (Vitamin D analogue), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0) Total (95% CI) 1489 974 100.0 % 0.0 [-0.01, 0.01] Total events: 0 (Vitamin D analogue), 0 (Placebo) Heterogeneity: Tau² = 0.0; Chi² = 0.0, df = 15 (P = 1.00); $I^2 = 0.0\%$ Test for overall effect: Z = 0.0 (P = 1.0) Test for subgroup differences: $Chi^2 = 0.0$, df = 6 (P = 1.00), $I^2 = 0.0\%$

> - I -0.5 0 0.5 I Favours vitamin D analogue Favours placebo

(... Continued)

Analysis 2.1. Comparison 2 Corticosteroid (potent) versus placebo, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: I IAGI

Study or subgroup	Corticosteroi (potent) N	d Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
l Betamethasone dipropiona	te OD	()		()			
Fleming 2010 (P)	78	2.03 (1.06)	28	2.64 (0.62)	-	8.8 %	-0.63 [-1.07, -0.19]
Kaufmann 2002 (P)	476	1.8992 (0.94)	157	2.68 (0.89)	-	15.4 %	-0.84 [-1.03, -0.65]
Subtotal (95% CI)	554		185		•	24.2 %	-0.81 [-0.98, -0.64]
Heterogeneity: $Tau^2 = 0.0$; C		f = 1 (P = 0.38);				_ 11_ /0	
Test for overall effect: $Z = 9.2$		· /					
2 Betamethasone dipropiona	te twice daily	,					
Papp 2003 (P)	312	-3.44 (1.05)	107	-1.92 (1.07)	-	3.9 %	-1.44 [-1.68, -1.20]
Vanderploeg 1976	17	-3.24 (0.97)	16	-2.06 (0.93)	-	4.4 %	-1.21 [-1.96, -0.46]
Wortzel 1975 (1)	39	-2.97 (1.09)	37	-1.7 (1.15)	-	7.9 %	-1.12 [-1.61, -0.64]
Wortzel 1975 (2)	5	-2.2 (1.1)	4	-1.25 (0.5)		1.4 %	-0.94 [-2.38, 0.49]
Subtotal (95% CI) Heterogeneity: Tau ² = 0.0; C	373 hi² = 1.79, dt	f = 3 (P = 0.62);	164 ² =0.0%		•	27.6 %	-1.35 [-1.56, -1.15]
Test for overall effect: $Z = 12$		0001)					
3 Betamethasone dipropiona	te, maintenar	nce					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	2						
Test for overall effect: not ap	olicable						
4 Betamethasone valerate							
Stein 2001	37	-3.1 (1.195)	37	-1.4 (1.195)	-	7.4 %	-1.41 [-1.92, -0.90]
Subtotal (95% CI)	37		37		•	7.4 %	-1.41 [-1.92, -0.90]
Heterogeneity: not applicable	2						
Test for overall effect: $Z = 5.3$	39 (P < 0.000	01)					
5 Budesonide							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable							
Test for overall effect: not ap	olicable						
6 Desonide Greenspan 1993	56	-2.8 (0.98)	20	-2 (0.98)	-	7.1 %	-0.81 [-1.34, -0.28]
		2.0 (0.70)		2 (0.70)			2
Subtotal (95% CI)	56		20		•	7.1 %	-0.81 [-1.34, -0.28]
Heterogeneity: not applicable Test for overall effect: $Z = 3.0$		ודי					
Test for overall effect: $Z = 3.0$	JU (F – 0.002	-/)					
				-10) -5 0 5 1	0	
				- ro Favours co			

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Study or subgroup	Corticosteroid (potent)		Placebo		Std. Mean Difference	Weight	(Continued Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
7 Diflorasone diacetate							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicat	ble						
Test for overall effect: not a	pplicable						
8 Fluticasone propionate							
Olsen 1996 (1)	88	-2.9 (1.37)	90	-1.7 (1.18)	-	11.9 %	-0.94 [-1.25, -0.63]
Olsen 1996 (2)	105	-2.8 (1.22)	100	-1.7 (1.15)	-	12.5 %	-0.92 [-1.21, -0.64]
Subtotal (95% CI)	193		190		•	24.4 %	-0.93 [-1.14, -0.72]
Heterogeneity: Tau ² = 0.0; Test for overall effect: Z = 8 9 Hydrocortisone buteprat Subtotal (95% CI) Heterogeneity: not applicab Test for overall effect: not a	3.63 (P < 0.0000 e 0	· ,	0.0%				Not estimable
10 Mometasone furoate Medansky 1987	50	-1.96 (0.9)	45	-1.24 (1)	-	9.3 %	-0.75 [-1.17, -0.34]
,		-1.70 (0.7)		-1.24 (1)			
Subtotal (95% CI) Heterogeneity: not applicab Test for overall effect: Z = 3		1)	45		•	9.3 %	-0.75 [-1.17, -0.34]
Total (95% CI)	1263	•)	641		•	100.0 %	-1.00 [-1.18, -0.82]
Heterogeneity: Tau ² = 0.05 Test for overall effect: Z =	; $Chi^2 = 23.60$, d 10.87 (P < 0.0000		,				

Favours corticosteroid Favours placebo

Analysis 2.2. Comparison 2 Corticosteroid (potent) versus placebo, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: 2 TSS

St Mea Difference	Weight	Std. Mean Difference		Placebo		Corticosteroid (potent)	Co Study or subgroup
IV,Random,95% (0	IV,Random,95% CI	Mean(SD)	Ν	Mean(SD)	N	,
						ite OD	I Betamethasone dipropionate
-0.74 [-1.16, -0.32	15.4 %	=	6.6 (2.8)	47	4.5 (2.8)	46	Lane 1983
-0.74 [-1.16, -0.32	15.4 %	•		47		46	Subtotal (95% CI) Heterogeneity: not applicable
					3)		Test for overall effect: $Z = 3.46$
						te twice daily	2 Betamethasone dipropionate
-0.77 [-1.48, -0.06	8.3 %	-8-	5.4 (2.8)	16	3.2 (2.8)	17	Vanderploeg 1976
-0.77 [-1.48, -0.06	8.3 %	•		16		17	Subtotal (95% CI)
						9	Heterogeneity: not applicable
						II (P = 0.034)	Test for overall effect: $Z = 2.1$
					9	ite, maintenance	3 Betamethasone dipropionate
-0.95 [-1.62, -0.27	8.9 %	-	2.12 (1.33)	19	0.83 (1.33)	19	Katz 1987a
-0.95 [-1.62, -0.27	8.9 %	•		19		19	Subtotal (95% CI)
						5	Heterogeneity: not applicable
)	76 (P = 0.0058)	Test for overall effect: $Z = 2.76$
							4 Betamethasone valerate
-1.09 [-2.00, -0.18	5.7 %	-	7.75 (2.45)	11	4.92 (2.53)		Ormerod 1997
-1.09 [-2.00, -0.18	5.7 %	•		11		11	Subtotal (95% CI)
						9	Heterogeneity: not applicable
						36 (P = 0.018)	Test for overall effect: $Z = 2.36$
				-			5 Budesonide
Not estimabl				0		0	Subtotal (95% CI)
							Heterogeneity: not applicable
						plicable	Test for overall effect: not appl 6 Desonide
-1.16 [-1.70, -0.61	11.7 %		4.11 (0.65)	20	3.07 (0.96)	56	Greenspan 1993
-1.16 [-1.70, -0.61	11.7 %	•	. /	20	. ,	56	Subtotal (95% CI)
1.10 [1.7 0, 0.01	11./ /0			20			Heterogeneity: not applicable
					32)		Test for overall effect: $Z = 4.16$
					,		7 Diflorasone diacetate
-0.32 [-0.73, 0.09	15.8 %	-	6.6 (2.8)	47	5.7 (2.8)	46	Lane 1983
-0.32 [-0.73, 0.09	15.8 %	•		47		46	Subtotal (95% CI)

Favours corticosteroid

Favours placebo

(Continued . . .)

Study or subgroup	Corticosteroid (potent)		Placebo		Std. Mean Difference	Weight	(Continued) Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
Heterogeneity: not applicabl							
Test for overall effect: $Z = I$.53 (P = 0.13)						
8 Fluticasone propionate							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicabl	e						
Test for overall effect: not ap	plicable						
9 Hydrocortisone buteprate	2						
Sears 1997	78	-2 (1.69)	83	-1.3 (1.32)	•	19.3 %	-0.46 [-0.77, -0.15]
Subtotal (95% CI)	78		83		•	19.3 %	-0.46 [-0.77, -0.15]
Heterogeneity: not applicabl	e						
Test for overall effect: $Z = 2$.89 (P = 0.0039)						
10 Mometasone furoate							
Medansky 1987	50	2.7 (1.33)	45	4.2 (1.33)	-	14.9 %	-1.12 [-1.55, -0.68]
Subtotal (95% CI)	50		45		•	14.9 %	-1.12 [-1.55, -0.68]
Heterogeneity: not applicabl	e						
Test for overall effect: $Z = 5$.05 (P < 0.0000))					
Total (95% CI)	323		288		•	100.0 %	-0.77 [-1.01, -0.52]
Heterogeneity: Tau ² = 0.05;	$Chi^2 = 13.13, dt$	f = 7 (P = 0.07);	; l ² =47%				
Test for overall effect: $Z = 6$.19 (P < 0.0000))					
Test for subgroup difference			7), $ ^2 = 47\%$				

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10

Favours placebo

Favours corticosteroid

Analysis 2.3. Comparison 2 Corticosteroid (potent) versus placebo, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: 3 PASI

Study or subgroup	Corticosteroid (potent) N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95%	Weight	Std. Mean Difference IV,Random,95% CI
l Betamethasone dipropiona	ate OD						
Fleming 2010 (P)	78	3.9 (3.8)	28	5.5 (3.5)	-	25.5 %	-0.43 [-0.86, 0.01]
Kaufmann 2002 (P)	476	-0.57 (0.3)	157	-0.23 (0.34)		38.4 %	-1.09 [-1.28, -0.90]
Subtotal (95% CI)	554	. ,	185		•	63.9 %	-0.79 [-1.44, -0.14]
Heterogeneity: $Tau^2 = 0.19$;		f = 1 (P = 0.01)	-			03.7 /0	-0./) [-1.44, -0.14]
Test for overall effect: $Z = 2$.		,	,1 -0770				
2 Betamethasone dipropiona	· · · ·						
Papp 2003 (P)	312	-0.63 (0.27)	107	-0.29 (0.31)		36.1 %	-1.21 [-1.44, -0.97]
SL+-+-1 (050/ CI)	210	. ,	107		•	26.1.0/	
Subtotal (95% CI)	312		107		•	36.1 %	-1.21 [-1.44, -0.97]
Heterogeneity: not applicable Test for overall effect: $Z = 10$							
	`	,					
3 Betamethasone dipropiona Subtotal (95% CI)	ate, maintenanc 0	e	0				Not estimable
Heterogeneity: not applicable	-		U				Not estimable
Test for overall effect: not ap							
4 Betamethasone valerate	plicable						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	e						
Test for overall effect: not ap							
5 Budesonide							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	e						
Test for overall effect: not ap	plicable						
6 Desonide							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	e						
Test for overall effect: not ap	plicable						
7 Diflorasone diacetate							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable							
Test for overall effect: not ap	plicable						
8 Fluticasone propionate	~		~				.
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	e						
						I	
						5 10	
				Favours c	orticosteroid Favo	ours placebo	(Continued)
							(Continued,

Study or subgroup	Corticosteroid (potent)	Placebo		Std. Mean Difference	Weight	(Continued) Std. Mean Difference
	Ν	Mean(SD) N	Mean(SD)	IV,Random,95% Cl		IV,Random,95% CI
Test for overall effect: not	t applicable					
9 Hydrocortisone butepr	ate					
Subtotal (95% CI)	0	0				Not estimable
Heterogeneity: not applic	able					
Test for overall effect: not	t applicable					
10 Mometasone furoate						
Subtotal (95% CI)	0	0				Not estimable
Heterogeneity: not applic	able					
Test for overall effect: not	t applicable					
Total (95% CI)	866	292		•	100.0 %	-0.97 [-1.31, -0.62]
Heterogeneity: $Tau^2 = 0.0$	07; Chi ² = 9.79, df	= 2 (P = 0.01); I ² =80%				
Test for overall effect: Z =	= 5.54 (P < 0.0000	I)				
Test for subgroup differer	nces: Chi ² = 1.40, c	If = 1 (P = 0.24), $I^2 = 29\%$				
			-	0 -5 0 5	10	

Favours corticosteroid Favours placebo

Analysis 2.5. Comparison 2 Corticosteroid (potent) versus placebo, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Corticosteroid (potent) N	d Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV.Random,95% Cl	Weight	Std. Mean Difference IV.Random,95% CI
		1 leal1(3D)	IN	Triedi (SD)			TV,IVandonii,75% Ci
I Betamethasone dipropion Fleming 2010 (P)	ate OD 78	2.03 (1.06)	28	2.64 (0.62)	-	6.5 %	-0.63 [-1.07, -0.19]
0 ()		· · · ·		× /			
Kaufmann 2002 (P)	476	1.8992 (0.94)	157	2.68 (0.89)	•	9.9 %	-0.84 [-1.03, -0.65]
Lane 1983	46	4.5 (2.8)	47	6.6 (2.8)	-	6.7 %	-0.74 [-1.16, -0.32]
Subtotal (95% CI)	600		232		•	23.2 %	-0.80 [-0.96, -0.64]
Heterogeneity: $Tau^2 = 0.0$; ($Chi^2 = 0.84, df$	= 2 (P = 0.66);	l ² =0.0%				
Test for overall effect: $Z = 9$	9.86 (P < 0.000	01)					
2 Betamethasone dipropion	ate twice daily						
Papp 2003 (P)	312	-3.44 (1.05)	107	-1.92 (1.07)	•	9.2 %	-1.44 [-1.68, -1.20]
Vanderploeg 1976	17	-3.24 (0.97)	16	-2.06 (0.93)		3.6 %	-1.21 [-1.96, -0.46]
Wortzel 1975 (1)	39	-2.97 (1.09)	37	-1.7 (1.15)	•	6.0 %	-1.12 [-1.61, -0.64]
Wortzel 1975 (2)	5	-2.2 (.)	4	-1.25 (0.5)		1.3 %	-0.94 [-2.38, 0.49]
Subtotal (95% CI)	373		164		•	20.1 %	-1.35 [-1.56, -1.15]
Heterogeneity: Tau ² = 0.0; 6 Test for overall effect: Z = 1 3 Betamethasone dipropion Katz 1987a	2.96 (P < 0.00	001)	² =0.0%	2.12 (1.33)	+	4.2 %	-0.95 [-1.62, -0.27]
Subtotal (95% CI)	19		19			4.2 %	-0.95 [-1.62, -0.27]
Heterogeneity: not applicab Test for overall effect: Z = 2 4 Betamethasone valerate Ormerod 1997	le	8) 4.92 (2.53)		7.75 (2.45)	_	4.2 70	-1.09 [-2.00, -0.18]
Stein 2001	37	-3.1 (1.195)	37	-1.4 (1.195)	-	5.7 %	-1.41 [-1.92, -0.90]
Subtotal (95% CI)	48		48		•	8.4 %	-1.33 [-1.78, -0.89]
Heterogeneity: Tau ² = 0.0; 0 Test for overall effect: $Z = 5$ 5 Budesonide	Chi ² = 0.35, df	. ,					100 [100, 000]
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicab	le						
Test for overall effect: not ap	oplicable						
					10 -5 0 5	10	
				Favours	corticosteroid Favours pl	acebo	
							(Continued)

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(... Continued) Std. Std. Corticosteroid Mean Mean Difference Difference Study or subgroup Placebo (potent) Weight IV,Random,95% CI Ν Mean(SD) Ν Mean(SD) IV,Random,95% CI 6 Desonide Greenspan 1993 . 56 -2.8 (0.98) 20 -2 (0.98) 5.5 % -0.81 [-1.34, -0.28] Subtotal (95% CI) -0.81 [-1.34, -0.28] 56 20 5.5 % Heterogeneity: not applicable Test for overall effect: Z = 3.00 (P = 0.0027) 7 Diflorasone diacetate Lane 1983 46 5.7 (2.8) 47 6.6 (2.8) 6.9 % -0.32 [-0.73, 0.09] Subtotal (95% CI) 6.9 % -0.32 [-0.73, 0.09] 46 47 Heterogeneity: not applicable Test for overall effect: Z = 1.53 (P = 0.13) 8 Fluticasone propionate Olsen 1996 (1) 88 -2.9 (1.37) 90 -1.7 (1.18) . 8.2 % -0.94 [-1.25, -0.63] Olsen 1996 (2) 105 -2.8 (1.22) 100 -1.7 (1.15) 8.6 % -0.92 [-1.21, -0.64] Subtotal (95% CI) -0.93 [-1.14, -0.72] 193 190 16.8 % Heterogeneity: Tau² = 0.0; Chi² = 0.00, df = 1 (P = 0.96); $I^2 = 0.0\%$ Test for overall effect: Z = 8.63 (P < 0.00001) 9 Hydrocortisone buteprate Sears 1997 78 -2 (1.69) 83 -1.3 (1.32) 8.2 % -0.46 [-0.77, -0.15] Subtotal (95% CI) 78 83 8.2 % -0.46 [-0.77, -0.15] Heterogeneity: not applicable Test for overall effect: Z = 2.89 (P = 0.0039) 10 Mometasone furoate Medansky 1987 50 6.8 % -0.75 [-1.17, -0.34] -1.96 (0.9) 45 -1.24 (1) Subtotal (95% CI) 50 45 6.8 % -0.75 [-1.17, -0.34] Heterogeneity: not applicable Test for overall effect: Z = 3.54 (P = 0.00041) Total (95% CI) 848 100.0 % -0.89 [-1.06, -0.72] 1463 Heterogeneity: Tau² = 0.07; Chi² = 43.03, df = 15 (P = 0.00016); $I^2 = 65\%$ Test for overall effect: Z = 10.07 (P < 0.00001) Test for subgroup differences: $Chi^2 = 40.04$, df = 8 (P = 0.00), $I^2 = 80\%$

-10 -5 0

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Favours corticosteroid Favours placebo

Analysis 2.6. Comparison 2 Corticosteroid (potent) versus placebo, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: 6 Total withdrawals

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference M- H,Random,95%	Weight	Risk Difference M- H,Random,9
	n/N	n/N	Cl		CI
I Betamethasone dipropionate (Fleming 2010 (P)	OD 5/83	12/40	-	10.6 %	-0.24 [-0.39, -0.09]
Kaufmann 2002 (P)	22/476	25/157	*	14.8 %	-0.11 [-0.17, -0.05]
Subtotal (95% CI)	559	197	•	25.4 %	-0.16 [-0.28, -0.04]
Total events: 27 (Corticosteroid Heterogeneity: Tau ² = 0.00; Chi Test for overall effect: $Z = 2.58$ (² = 2.38, df = 1 (P =	,			
2 Betamethasone dipropionate t	,				
Papp 2003 (P)	17/313	12/108	-	14.6 %	-0.06 [-0.12, 0.01]
Subtotal (95% CI)	313	108	•	14.6 %	-0.06 [-0.12, 0.01]
Total events: 17 (Corticosteroid Heterogeneity: not applicable Test for overall effect: Z = 1.73 (3 Betamethasone dipropionate,	(P = 0.084)	00)			
Katz 1987a	6/20	16/20	_ 	6.2 %	-0.50 [-0.77, -0.23]
Katz 1991a	16/48	35/46		9.3 %	-0.43 [-0.61, -0.25]
Subtotal (95% CI)	68	66	•	15.5 %	-0.45 [-0.60, -0.30]
Total events: 22 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 5.88 (4 Betamethasone valerate Stein 2001	= 0.19, df = 1 (P =	,		12.3 %	0.0 [-0.12, 0.12
Subtotal (95% CI)	40	40	•	12.3 %	0.0 [-0.12, 0.12]
Total events: 3 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (F 5 Budesonide	(potent)), 3 (Placebo			12.9 70	0.0 [-0.12, 0.12]
Agrup 1981	0/11	0/11	+	10.2 %	0.0 [-0.16, 0.16]
0 1	11	11	+	10.2 %	0.0 [-0.16, 0.16]

(Continued . . .)

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference M-	Weight	(Continued Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
6 Desonide					
Greenspan 1993	4/60	5/20		8.5 %	-0.18 [-0.38, 0.02]
Subtotal (95% CI)	60	20	•	8.5 %	-0.18 [-0.38, 0.02]
Total events: 4 (Corticosteroi	d (potent)), 5 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 1.8	0 (P = 0.072)				
7 Diflorasone diacetate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroi	d (potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not app	licable				
8 Fluticasone propionate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroi	d (potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not app	licable				
9 Hydrocortisone buteprate					
Sears 1997	10/84	11/96	-	13.4 %	0.00 [-0.09, 0.10]
Subtotal (95% CI)	84	96	+	13.4 %	0.00 [-0.09, 0.10]
Total events: 10 (Corticosterc	oid (potent)), II (Placebo	o)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	9 (P = 0.93)				
10 Mometasone furoate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroi	d (potent)), 0 (Placebo)				
Heterogeneity: not applicable	u ,, (
Test for overall effect: not app	licable				
Total (95% CI)	1135	538	•	100.0 %	-0.14 [-0.22, -0.05]
Total events: 83 (Corticosterc	oid (potent)), 119 (Placel	00)			
Heterogeneity: $Tau^2 = 0.01$; C	Chi² = 38.78, df = 8 (P<	0.0000 l); l ² =79%			
Test for overall effect: $Z = 3.1$					
Test for subgroup differences:	Chi ² = 32.44, df = 6 (P	= 0.00), I ² =82%			
- '	· · · · · · · · · · · · · · · · · · ·				

Favours corticosteroid Favours placebo

Analysis 2.7. Comparison 2 Corticosteroid (potent) versus placebo, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference M- H,Random,95%	Weight	Risk Difference M- H,Random,
	n/N	n/N	Cl		C
Betamethasone dipropionate	OD				
Kaufmann 2002 (P)	5/476	12/157	-	16.9 %	-0.07 [-0.11, -0.02]
Subtotal (95% CI)	476	157	•	16.9 %	-0.07 [-0.11, -0.02]
Total events: 5 (Corticosteroid ((potent)), 12 (Placebo)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 3.04$	(P = 0.0024)				
2 Betamethasone dipropionate	twice daily				
Vanderploeg 1976	0/17	0/16		6.9 %	0.0 [-0.11, 0.11]
Subtotal (95% CI)	17	16	+	6.9 %	0.0 [-0.11, 0.11]
Total events: 0 (Corticosteroid ((potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ (F	P = 1.0)				
3 Betamethasone dipropionate,	maintenance				
Katz 1987a	0/20	0/20	-	8.8 %	0.0 [-0.09, 0.09
Katz 1991a	0/48	0/46	-	17.3 %	0.0 [-0.04, 0.04
Subtotal (95% CI)	68	66	•	26.0 %	0.0 [-0.04, 0.04]
Fotal events: 0 (Corticosteroid ((potent)), 0 (Placebo)				
Heterogeneity: Tau ² = 0.0; Chi ²	= 0.0, df = 1 (P = 1.	00); l ² =0.0%			
Test for overall effect: $Z = 0.0$ (F					
1 Betamethasone valerate					
Stein 2001	3/40	0/40	-	8.8 %	0.08 [-0.02, 0.17]
Subtotal (95% CI)	40	40	•	8.8 %	0.08 [-0.02, 0.17]
Total events: 3 (Corticosteroid ((potent)), 0 (Placebo)				
Heterogeneity: not applicable	. ,, , ,				
Test for overall effect: $Z = 1.60$	(P = 0.11)				
5 Budesonide	. ,				
Agrup 1981	0/11	0/11		4.0 %	0.0 [-0.16, 0.16
Subtotal (95% CI)	11	11	•	4.0 %	0.0 [-0.16, 0.16]
Fotal events: 0 (Corticosteroid ((potent)), 0 (Placebo)				_ ,
Heterogeneity: not applicable	、 /				
e , , , , ,	P = 1.0)				
Test for overall effect: $Z = 0.0$ (F					

(Continued ...)

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference M- H,Random,95%	Weight	(Continuec Risk Difference M- H,Random,9
	n/N	n/N	Cl		Cl
6 Desonide					
Greenspan 1993	0/60	2/20		4.9 %	-0.10 [-0.24, 0.04]
Subtotal (95% CI)	60	20	•	4.9 %	-0.10 [-0.24, 0.04]
Total events: 0 (Corticosteroi	d (potent)), 2 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 1.4$	10 (P = 0.16)				
7 Diflorasone diacetate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroi	d (potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not app	olicable				
8 Fluticasone propionate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroi	d (potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not app	olicable				
9 Hydrocortisone buteprate					
Sears 1997	1/94	0/96	•	19.5 %	0.01 [-0.02, 0.04]
Subtotal (95% CI)	94	96	•	19.5 %	0.01 [-0.02, 0.04]
Total events: (Corticosteroi	d (potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.7$	72 (P = 0.47)				
10 Mometasone furoate					
Medansky 1987	0/61	3/59	-	13.0 %	-0.05 [-0.11, 0.01]
,					
Subtotal (95% CI)	61	59	•	13.0 %	-0.05 [-0.11, 0.01]
Total events: 0 (Corticosteroi	. ,, , , ,				
Heterogeneity: not applicable					
Test for overall effect: $Z = 1.5$	· · · · ·			100.0.0/	
Total (95% CI)	827	465	Ţ	100.0 %	-0.01 [-0.05, 0.02]
Total events: 9 (Corticosteroi	u // (,			
Heterogeneity: $Tau^2 = 0.00$; (= 0.01); 14 =61%			
Test for overall effect: $Z = 0.7$	· · · · ·	0.00) 12 5.464			
Test for subgroup differences:	$Chr^2 = 15.38, df = 7 (P$	= 0.03), I ² =54%			

Favours corticosteroid (potent) Favours placebo

Analysis 2.8. Comparison 2 Corticosteroid (potent) versus placebo, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference	Weight	Risk Difference
	n/N	n/N	M- H,Random,95% Cl		M- H,Random,9 CI
I Betamethasone dipropionate	OD				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid (
Heterogeneity: not applicable	(======)), = (: :======)				
Test for overall effect: not applic	able				
2 Betamethasone dipropionate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid (-	-			1.0000000000000000000000000000000000000
Heterogeneity: not applicable	(potenty); o (nacebo)	,			
Test for overall effect: not applic	able				
3 Betamethasone dipropionate,					
Katz 1987a	5/20	15/20		31.5 %	-0.50 [-0.77, -0.23]
Katz 1707a	5/20	15/20		51.5 %	-0.50 [-0.77, -0.25]
Katz 1991a	16/46	35/44		68.5 %	-0.45 [-0.63, -0.27]
Subtotal (95% CI)	66	64	•	100.0 %	-0.46 [-0.61, -0.31]
. ,			•	100.0 %	-0.46 [-0.61, -0.31]
Total events: 21 (Corticosteroid	(potent)), 50 (Placeb	00)	•	100.0 %	-0.46 [-0.61, -0.31]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ²	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0	00)	•	100.0 %	-0.46 [-0.61, -0.31]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0	00)	•	100.0 %	-0.46 [-0.61, -0.31]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: $Z = 6.04$ 4 Betamethasone valerate	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001)	0.75); I ² =0.0%	•		
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001	l (potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40	0,75); I ² =0.0%	•	100.0 %	0.0 [-0.05, 0.05]
Subtotal (95% CI)	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40	00) 0.75); I ² =0.0% 0/40 40	-		0.0 [-0.05, 0.05]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI)	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40	00) 0.75); I ² =0.0% 0/40 40	•	100.0 %	-0.46 [-0.61, -0.31] 0.0 [-0.05, 0.05] 0.0 [-0.05, 0.05]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40	00) 0.75); I ² =0.0% 0/40 40	•	100.0 %	0.0 [-0.05, 0.05]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable	l (potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo)	00) 0.75); I ² =0.0% 0/40 40	•	100.0 %	0.0 [-0.05, 0.05]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f	l (potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo)	00) 0.75); I ² =0.0% 0/40 40	•	100.0 %	0.0 [-0.05, 0.05]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f	l (potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo)	00) 0.75); I ² =0.0% 0/40 40	•	100.0 %	0.0 [-0.05, 0.05] 0.0 [-0.05, 0.05]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 5 Budesonide Agrup 1981	I (potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo) P = 1.0)	00) 0.75); I ² =0.0% 0/40 40	•	100.0 % 100.0 %	0.0 [-0.05, 0.05] 0.0 [-0.05, 0.05] 0.0 [-0.16, 0.16]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 5 Budesonide Agrup 1981 Subtotal (95% CI)	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo) P = 1.0) 0/11 11	00) 0.75); I ² =0.0% 0/40 40 0/11 11	•	100.0 % 100.0 %	0.0 [-0.05, 0.05] 0.0 [-0.05, 0.05] 0.0 [-0.16, 0.16]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 5 Budesonide	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo) P = 1.0) 0/11 11	00) 0.75); I ² =0.0% 0/40 40 0/11 11	•	100.0 % 100.0 %	0.0 [-0.05, 0.05] 0.0 [-0.05, 0.05] 0.0 [-0.16, 0.16]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 5 Budesonide Agrup 1981 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo) P = 1.0) 0/11 11 (potent)), 0 (Placebo)	00) 0.75); I ² =0.0% 0/40 40 0/11 11	•	100.0 % 100.0 %	0.0 [-0.05, 0.05] 0.0 [-0.05, 0.05] 0.0 [-0.16, 0.16]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 5 Budesonide Agrup 1981 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo) P = 1.0) 0/11 11 (potent)), 0 (Placebo)	00) 0.75); I ² =0.0% 0/40 40 0/11 11	•	100.0 % 100.0 %	0.0 [-0.05, 0.05] 0.0 [-0.05, 0.05] 0.0 [-0.16, 0.16]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 5 Budesonide Agrup 1981 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo) P = 1.0) 0/11 11 (potent)), 0 (Placebo)	00) 0.75); I ² =0.0% 0/40 40 0/11 11		100.0 % 100.0 %	0.0 [-0.05, 0.05] 0.0 [-0.05, 0.05] 0.0 [-0.16, 0.16] 0.0 [-0.16, 0.16]
Total events: 21 (Corticosteroid Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 6.04 4 Betamethasone valerate Stein 2001 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 5 Budesonide Agrup 1981 Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 6 Desonide	(potent)), 50 (Placeb = 0.10, df = 1 (P = 0 (P < 0.00001) 0/40 40 (potent)), 0 (Placebo) P = 1.0) 0/11 11 (potent)), 0 (Placebo) P = 1.0)	00) 0.75); 1 ² =0.0% 0/40 40 0/1 1 11		100.0 % 100.0 % 100.0 % 100.0 %	0.0 [-0.05, 0.05]

(Continued . . .)

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference M- H,Random <u>,</u> 95%	Weight	(Continued) Risk Difference M- H,Random <u>,</u> 95%
	n/N	n/N	Cl		CI
Total events: 0 (Corticosteroid	(potent)), 0 (Placebo)				
Heterogeneity: not applicable Test for overall effect: $Z = 0.0$ ((D - 1.0)				
7 Diflorasone diacetate	(P = 1.0)				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid	-	0			Not estimable
Heterogeneity: not applicable	(potent)), o (nacebo)				
Test for overall effect: not appli	cable				
8 Fluticasone propionate	Cable				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid	-	Ū			The commande
Heterogeneity: not applicable	(potent)), o (nacebo)				
Test for overall effect: not appli	cable				
9 Hydrocortisone buteprate					
Sears 1997	0/94	0/96		100.0 %	0.0 [-0.02, 0.02]
Subtotal (95% CI)	94	96	•	100.0 %	0.0 [-0.02, 0.02]
Total events: 0 (Corticosteroid		<i>)</i> 0		100.0 /0	0.0 [-0.02, 0.02]
Heterogeneity: not applicable	(potent)), o (nacebo)				
Test for overall effect: $Z = 0.0$ ((P = 1.0)				
	(1.0)				
10 Mometasone furoate	0	0			Not estimable
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid	(potent)), 0 (Placebo)				
Heterogeneity: not applicable Test for overall effect: not appli	eable				
Test for subgroup differences: C		- 0.00) 12 - 000/			
rest for subgroup differences: C	∠iii — 33.74, úi — 4 (P	− 0.00), I ⁻ −07%			
			-I -0.5 0 0.5 I		
		Favours corticoste	eroid (potent) Favours placebo)	

Analysis 2.9. Comparison 2 Corticosteroid (potent) versus placebo, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: 9 Adverse events (local)

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
I Betamethasone dipropionate	e OD				
Fleming 2010 (P)	7/83	10/40		5.1 %	-0.17 [-0.31, -0.02]
Kaufmann 2002 (P)	23/476	21/157	+	14.1 %	-0.09 [-0.14, -0.03]
Subtotal (95% CI)	559	197	•	19.2 %	-0.10 [-0.15, -0.04]
Total events: 30 (Corticostero Heterogeneity: Tau ² = 0.00; C Test for overall effect: $Z = 3.42$	$hi^2 = 1.03, df = 1 (P = 3 (P = 0.00060)$				
2 Betamethasone dipropionate Papp 2003 (P)	e twice daily 27/313	17/108	-	11.4 %	-0.07 [-0.15, 0.00]
Vanderploeg 1976	0/17	0/16	-	7.5 %	0.0 [-0.11, 0.11]
Subtotal (95% CI)	330	124	•	18.9 %	-0.05 [-0.12, 0.03]
Test for overall effect: $Z = 1.19$	9 (P = 0.23)				
0 ,					
Test for overall effect: $Z = 1.19$ 3 Betamethasone dipropionate	9 (P = 0.23) e, maintenance	0/20	+	Q 3 %	
Test for overall effect: Z = 1.19 3 Betamethasone dipropionate Katz 1987a	9 (P = 0.23) e, maintenance 0/20	0/20	-	9.3 %	-
Test for overall effect: $Z = 1.19$ Betamethasone dipropionate	9 (P = 0.23) e, maintenance	0/20 0/46	+	9.3 %	-
Test for overall effect: Z = 1.19 3 Betamethasone dipropionate Katz 1987a Katz 1991a Subtotal (95% CI)	9 (P = 0.23) e, maintenance 0/20 0/48 68	0/46 66	-		0.0 [-0.04, 0.04]
Test for overall effect: Z = 1.19 3 Betamethasone dipropionate Katz 1987a Katz 1991a Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0	9 (P = 0.23) e, maintenance 0/20 0/48 68 (potent)), 0 (Placebo) i ² = 0.0, df = 1 (P = 1.	0/46 66	•	16.6 %	0.0 [-0.04, 0.04]
Test for overall effect: Z = 1.19 3 Betamethasone dipropionate Katz 1987a Katz 1991a Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0 4 Betamethasone valerate	9 (P = 0.23) e, maintenance 0/20 0/48 68 6 (potent)), 0 (Placebo) i ² = 0.0, df = 1 (P = 1. (P = 1.0)	0/46 66 00); I ² =0.0%	•	16.6 %	0.0 [-0.04, 0.04]
Test for overall effect: Z = 1.19 3 Betamethasone dipropionate Katz 1987a Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0 4 Betamethasone valerate Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable	P (P = 0.23) e, maintenance 0/20 0/48 68 d (potent)), 0 (Placebo) i ² = 0.0, df = 1 (P = 1. (P = 1.0) 0 d (potent)), 0 (Placebo)	0/46 66 00); I ² =0.0% 0	•	16.6 %	0.0 [-0.09, 0.09] 0.0 [-0.04, 0.04] 0.0 [-0.04, 0.04] Not estimable
Test for overall effect: Z = 1.19 3 Betamethasone dipropionate Katz 1987a Katz 1991a Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0 4 Betamethasone valerate Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable Test for overall effect: not appli	P (P = 0.23) e, maintenance 0/20 0/48 68 d (potent)), 0 (Placebo) i ² = 0.0, df = 1 (P = 1. (P = 1.0) 0 d (potent)), 0 (Placebo)	0/46 66 00); I ² =0.0% 0	•	16.6 %	0.0 [-0.04, 0.04]
Test for overall effect: Z = 1.15 3 Betamethasone dipropionate Katz 1987a Katz 1991a Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0 4 Betamethasone valerate Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable Test for overall effect: not appl 5 Budesonide	9 (P = 0.23) e, maintenance 0/20 0/48 68 d (potent)), 0 (Placebo) i ² = 0.0, df = 1 (P = 1. (P = 1.0) 0 d (potent)), 0 (Placebo) iicable	0/46 66 00); I ² =0.0% 0	•	16.6 %	0.0 [-0.04, 0.04] 0.0 [-0.04, 0.04] Not estimable
Test for overall effect: Z = 1.15 3 Betamethasone dipropionate Katz 1987a Katz 1991a Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0 4 Betamethasone valerate Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable Test for overall effect: not appl 5 Budesonide Subtotal (95% CI)	P (P = 0.23) e, maintenance 0/20 0/48 68 d (potent)), 0 (Placebo) i ² = 0.0, df = 1 (P = 1. (P = 1.0) 0 d (potent)), 0 (Placebo) iicable 0	0/46 66 00); I ² =0.0% 0	•	16.6 %	0.0 [-0.04, 0.04]
Test for overall effect: Z = 1.15 3 Betamethasone dipropionate Katz 1987a Katz 1987a Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0 4 Betamethasone valerate Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable Test for overall effect: not appl 5 Budesonide Subtotal (95% CI) Total events: 0 (Corticosteroic	P (P = 0.23) e, maintenance 0/20 0/48 68 d (potent)), 0 (Placebo) i ² = 0.0, df = 1 (P = 1. (P = 1.0) 0 d (potent)), 0 (Placebo) iicable 0	0/46 66 00); I ² =0.0% 0	•	16.6 %	0.0 [-0.04, 0.04] 0.0 [-0.04, 0.04] Not estimable
	P (P = 0.23) e, maintenance 0/20 0/48 68 4 (potent)), 0 (Placebo) P = 1.0 0 4 (potent)), 0 (Placebo) 6 (potent)), 0 (Placebo) 6 (potent)), 0 (Placebo)	0/46 66 00); I ² =0.0% 0	•	16.6 %	0.0 [-0.04, 0.04] 0.0 [-0.04, 0.04] Not estimable
Test for overall effect: Z = 1.19 3 Betamethasone dipropionate Katz 1987a Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0 4 Betamethasone valerate Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable Test for overall effect: not appl 5 Budesonide Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable Total events: 0 (Corticosteroic Heterogeneity: not applicable	P (P = 0.23) e, maintenance 0/20 0/48 68 4 (potent)), 0 (Placebo) P = 1.0 0 4 (potent)), 0 (Placebo) 6 (potent)), 0 (Placebo) 6 (potent)), 0 (Placebo)	0/46 66 00); I ² =0.0% 0		16.6 %	0.0 [-0.04, 0.04 0.0 [-0.04, 0.04] Not estimable

(Continued . . .)

(... Continued) Risk Risk Corticosteroid Placebo Difference Difference Study or subgroup (potent) Weight M-M-H,Random,95% H,Random,95% n/N n/N ĆI Ć Greenspan 1993 3/60 1/20 7.6 % 0.0 [-0.11, 0.11] Subtotal (95% CI) 20 60 7.6% 0.0 [-0.11, 0.11] Total events: 3 (Corticosteroid (potent)), | (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0) 7 Diflorasone diacetate Subtotal (95% CI) 0 0 Not estimable Total events: 0 (Corticosteroid (potent)), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: not applicable 8 Fluticasone propionate Olsen 1996 (1) 13/193 12/190 15.2 % 0.00 [-0.05, 0.05] Subtotal (95% CI) 193 0.00 [-0.05, 0.05] 190 15.2 % Total events: 13 (Corticosteroid (potent)), 12 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.17 (P = 0.87) 9 Hydrocortisone buteprate Sears 1997 21/94 27/96 6.5 % -0.06 [-0.18, 0.07] Subtotal (95% CI) 94 96 -0.06 [-0.18, 0.07] 6.5 % Total events: 21 (Corticosteroid (potent)), 27 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.92 (P = 0.36) 10 Mometasone furoate Medansky 1987 5/61 11/59 6.7 % -0.10 [-0.23, 0.02] Subtotal (95% CI) 59 61 6.7 % -0.10 [-0.23, 0.02] Total events: 5 (Corticosteroid (potent)), 11 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 1.69 (P = 0.090) Total (95% CI) 1365 752 100.0 % -0.04 [-0.08, 0.00] Total events: 99 (Corticosteroid (potent)), 99 (Placebo) Heterogeneity: Tau² = 0.00; Chi² = 20.56, df = 9 (P = 0.01); $I^2 = 56\%$ Test for overall effect: Z = 1.98 (P = 0.047) Test for subgroup differences: $Chi^2 = 11.95$, df = 6 (P = 0.06), $I^2 = 50\%$ - | -0.5 0 0.5 I

Favours corticosteroid (potent) Favours placebo

Analysis 2.10. Comparison 2 Corticosteroid (potent) versus placebo, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 2 Corticosteroid (potent) versus placebo

Outcome: 10 Adverse events (systemic)

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference M- H,Random,95%	Weight	Risk Difference M- H,Random,95
	n/N	n/N	Cl		Cl
I Betamethasone dipropionate	OD				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid ((potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applic	able				
2 Betamethasone dipropionate	twice daily		\perp		
Papp 2003 (P)	0/313	0/108	-	81.6 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	313	108		81.6 %	0.0 [-0.01, 0.01]
Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f 3 Betamethasone dipropionate,	P = 1.0)				
Katz 1987a	0/20	1/20		0.9 %	-0.05 [-0.18, 0.08]
Katz 1991a	2/48	0/46		3.2 %	0.04 [-0.03, 0.11]
NdlZ 1771d	2/40	0/46		3.Z /o	0.04 [-0.05, 0.11]
Subtotal (95% CI)	68	66	+	4.1 %	0.01 [-0.07, 0.10]
Total events: 2 (Corticosteroid (Heterogeneity: Tau ² = 0.00; Chi Test for overall effect: $Z = 0.27$	(potent)), (Placebo) $i^2 = 1.59$, df = (P = 0		•	4.1 %	0.01 [-0.07, 0.10]
Subtotal (95% CI) Total events: 2 (Corticosteroid (Heterogeneity: Tau ² = 0.00; Chi Test for overall effect: Z = 0.27 4 Budesonide Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: not applic 5 Desonide	(potent)), (Placebo) i ² = 1.59, df = (P = 0 (P = 0.78) 0 (potent)), 0 (Placebo)		•	4.1 %	0.01 [-0.07, 0.10] Not estimable
Total events: 2 (Corticosteroid (Heterogeneity: Tau ² = 0.00; Chi Test for overall effect: Z = 0.27 4 Budesonide Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: not applic 5 Desonide	(potent)), (Placebo) i ² = 1.59, df = (P = 0 (P = 0.78) 0 (potent)), 0 (Placebo)	0.21); 1 ² =37%		4.1 %	
Total events: 2 (Corticosteroid (Heterogeneity: Tau ² = 0.00; Chi Test for overall effect: Z = 0.27 4 Budesonide Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: not applic 5 Desonide Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: not applicable	(potent)), (Placebo) i ² = 1.59, df = (P = 0 (P = 0.78) 0 (potent)), 0 (Placebo) table 0 (potent)), 0 (Placebo)	0.21); l ² =37%		4.1 %	Not estimable
Total events: 2 (Corticosteroid (Heterogeneity: Tau ² = 0.00; Chi Test for overall effect: Z = 0.27 4 Budesonide Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable Test for overall effect: not applic 5 Desonide Subtotal (95% CI) Total events: 0 (Corticosteroid (Heterogeneity: not applicable	(potent)), (Placebo) i ² = 1.59, df = (P = 0 (P = 0.78) 0 (potent)), 0 (Placebo) table 0 (potent)), 0 (Placebo)	0.21); l ² =37%		4.1 %	Not estimable

(Continued ...)

Study or subgroup	Corticosteroid (potent)	Placebo	Risk Difference M- H,Random,95%	Weight	(Continued) Risk Difference M- H,Random,95
	n/N	n/N	Cl		H,Random,23 Cl
Test for overall effect: not applie	cable				
7 Fluticasone propionate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid	(potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not appli	cable				
8 Hydrocortisone buteprate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid	(potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not appli	cable				
9 Betamethasone valerate					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Corticosteroid	(potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not appli	cable				
10 Mometasone furoate					
Medansky 1987	0/61	0/59	•	14.4 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	61	59	•	14.4 %	0.0 [-0.03, 0.03]
Total events: 0 (Corticosteroid	(potent)), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ ((P = 1.0)				
Total (95% CI)	442	233		100.0 %	0.00 [-0.01, 0.01]
Total events: 2 (Corticosteroid	(potent)), I (Placebo)				
Heterogeneity: Tau ² = 0.0; Chi ²	² = 2.16, df = 3 (P = 0	.54); l ² =0.0%			
Test for overall effect: $Z = 0.14$	(P = 0.89)				
Test for subgroup differences: C	$Chi^2 = 0.07, df = 2 (P = 1)$	= 0.96), I ² =0.0%			

-I -0.5 0 0.5 I

Favours corticosteroid (potent) Favours placebo

Analysis 3.1. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: I IAGI

Study or subgroup	Corticosteroid (very potent)		Placebo		Std. Mean Difference	Weight	Std. Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	IV,Random,95% C	3	IV,Random,95% CI
I Clobetasol propionate							
Beutner 2006	25	0.24 (0.44)	25	2.24 (1.05)	-	16.6 %	-2.45 [-3.19, -1.70]
Decroix 2004	189	-4.94 (1.08)	33	-2.23 (1.69)	-	21.9 %	-2.27 [-2.70, -1.85]
Jarratt 2006	60	1.1 (0.95)	60	2.72 (0.64)	-	21.7 %	-1.99 [-2.43, -1.55]
Lebwohl 2002	60	-2.85 (1.47)	19	-1.58 (1.17)	-	20.1 %	-0.89 [-1.43, -0.36]
Subtotal (95% CI)	334		137		•	80.4 %	-1.89 [-2.53, -1.24]
Heterogeneity: $Tau^2 = 0.36$; Chi ² = 18.67, d	= 3 (P = 0.0003	2); I ² =849	6			
Test for overall effect: $Z = $	5.74 (P < 0.0000)					
2 Halcinonide							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicat	ble						
Test for overall effect: not a	pplicable						
3 Halobetasol							
Bernhard 1991(2)	36	-2.97 (0.97)	33	-1.3 (0.85)	-	19.6 %	-1.81 [-2.37, -1.24]
Subtotal (95% CI)	36		33		•	19.6 %	-1.81 [-2.37, -1.24]
Heterogeneity: not applicat	ble						
Test for overall effect: $Z = $	6.26 (P < 0.0000))					
Total (95% CI)	370		170		•	100.0 %	-1.87 [-2.38, -1.36]
Heterogeneity: Tau ² = 0.26	; Chi ² = 18.76, dt	= 4 (P = 0.0008	8); I ² =799	6			
Test for overall effect: $Z = \frac{1}{2}$	7.21 (P < 0.0000))					
Test for subgroup difference	es: Chi² = 0.04, d	f = 1 (P = 0.85),	$ ^2 = 0.0\%$				
						L	
					-10 -5 0 5	10	
			F	Favours corticoster	roid (v potent) Favour	s placebo	

Analysis 3.2. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: 2 TSS

Study or subgroup	Corticosteroid (very potent) N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% (Weight	Std. Mean Difference IV,Random,95% CI
l Clobetasol propionate							
Gottlieb 2003	139	-4.01 (2.43)	140	-1.51 (2.43)		39.5 %	-1.03 [-1.28, -0.78]
Jorizzo 1997	35	-4.3 (1.98)	39	-1.6 (1.98)	-	28.7 %	-1.35 [-1.86, -0.84]
Lowe 2005	163	2.43 (2.03)	29	5.93 (1.85)	-	31.8 %	-1.74 [-2.17, -1.31]
Subtotal (95% CI)	337		208		•	100.0 %	-1.35 [-1.80, -0.89]
Heterogeneity: $Tau^2 = 0.12$; Chi ² = 8.10, df	= 2 (P = 0.02); I	² =75%				
Test for overall effect: $Z = $	5.83 (P < 0.0000)					
2 Halcinonide		, ,					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicab	le						
Test for overall effect: not a	pplicable						
3 Halobetasol							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicat	le						
Test for overall effect: not a	pplicable						
Total (95% CI)	337		208		•	100.0 %	-1.35 [-1.80, -0.89]
Heterogeneity: $Tau^2 = 0.12$; Chi ² = 8.10, df	= 2 (P = 0.02); I	² =75%				
Test for overall effect: $Z = S$	5.83 (P < 0.0000)					
Test for subgroup difference	es: Not applicable	1					
				-1	0 -5 0 5	10	
				Favours corticosteroi	id (v potent) Favour	s placebo	

Analysis 3.4. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: 4 PAGI

Stø Mea Differenc IV,Random,95% (Weight	Std. Mean Difference IV,Random,95% Cl	Mean(SD)	Placebo N	Mean(SD)	Corticosteroid (very potent) N	Study or subgroup
							Clobetasol propionate
-1.01 [-1.55, -0.47	13.4 %	-	-1.68 (1.25)	19	-3.07 (1.4)	60	Lebwohl 2002
-1.01 [-1.55, -0.47	13.4 %	•		19		60	Subtotal (95% CI)
						ble	Heterogeneity: not applica
)	3.65 (P = 0.00026	Test for overall effect: Z =
							2 Halcinonide
Not estimable				0		0	Subtotal (95% CI)
						ble	Heterogeneity: not applica
						applicable	Test for overall effect: not
							3 Halobetasol
-1.17 [-1.47, -0.86	41.7 %		-1.47 (1.29)	96	-2.92 (1.18)	96	Bernhard 1991 (1)
-1.33 [-1.62, -1.03	44.9 %	-	-0.72 (1.14)	108	-2.46 (1.45)	108	Katz 1991b
-1.25 [-1.46, -1.04	86.6 %	•		204		204	Subtotal (95% CI)
				=0.0%	I (P = 0.46); I ² :	; Chi ² = 0.55, df =	Heterogeneity: $Tau^2 = 0.0$;
					I)	II.54 (P < 0.0000	Test for overall effect: $Z =$
-1.22 [-1.42, -1.02	100.0 %	•		223		264	Total (95% CI)
				=0.0%	2 (P = 0.54); I ² :	; Chi ² = 1.23, df =	Heterogeneity: $Tau^2 = 0.0$;
					I)	12.08 (P < 0.0000	Test for overall effect: Z =
				2 =0.0%	F = (P = 0.4),	ces: $Chi^2 = 0.68$, df	Test for subgroup difference

Favours corticosteroid (v potent)

Favours placebo

Analysis 3.5. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Corticosteroid (very potent)		Placebo		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
Clobetasol propionate							
Beutner 2006	25	0.24 (0.44)	25	2.24 (1.05)		7.2 %	-2.45 [-3.19, -1.70]
Decroix 2004	189	-4.94 (1.08)	33	-2.23 (1.69)		10.3 %	-2.27 [-2.70, -1.85]
Gottlieb 2003	139	-4.01 (2.43)	140	-1.51 (2.43)	+	11.9 %	-1.03 [-1.28, -0.78]
Jarratt 2006	60	1.1 (0.95)	60	2.72 (0.64)		10.1 %	-1.99 [-2.43, -1.55]
Jorizzo 1997	35	-4.3 (1.98)	39	-1.6 (1.98)		9.4 %	-1.35 [-1.86, -0.84]
Lebwohl 2002	60	-2.85 (1.47)	19	-1.58 (1.17)		9.2 %	-0.89 [-1.43, -0.36]
Lowe 2005	163	2.43 (2.03)	29	5.93 (1.85)	-	10.2 %	-1.74 [-2.17, -1.31]
Subtotal (95% CI)	671		345		•	68.3 %	-1.65 [-2.10, -1.20]
2 Halcinonide Subtotal (95% CI)	0		0				Not estimable
	0		0				Not estimable
Heterogeneity: not applicab	le						
Test for overall effect: not a	pplicable						
3 Halobetasol Bernhard 1991 (1)	96	-2.92 (1.18)	96	-1.47 (1.29)	-	11.4 %	-1.17 [-1.47, -0.86]
		~ /		× /	_		
Bernhard 1991(2)	36	-2.97 (0.97)	33	-1.3 (0.85)		8.9 %	-1.81 [-2.37, -1.24]
Katz 1991b	108	-2.46 (1.45)	108	-0.72 (1.14)	-	11.5 %	-1.33 [-1.62, -1.03]
Subtotal (95% CI)	240		237		•	31.7 %	-1.36 [-1.65, -1.07]
Heterogeneity: $Tau^2 = 0.03$,	2 =47%				
Test for overall effect: $Z = 9$)					
Total (95% CI)	911		582		•	100.0 %	-1.56 [-1.87, -1.26]
			1, 12 -0.00/				
Heterogeneity: Tau ² = 0.18; Test for overall effect: $Z = 1$); = -02%				

-4 -2 0 2 4

Favours corticosteroid (v potent) Favours placebo

Analysis 3.6. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: 6 Total withdrawals

Corticosteroid (very potent)	Placebo	Risk Difference M- H Bandom 95%	Weight	Risk Difference M- H,Random,95
n/N	n/N	Cl		CI
2/27	2/27	-	9.1 %	0.0 [-0.14, 0.14]
5/189	4/33		11.1 %	-0.09 [-0.21, 0.02]
4/139	4/140	-	18.3 %	0.00 [-0.04, 0.04]
0/60	0/60	-	18.9 %	0.0 [-0.03, 0.03]
6/44	15/45		7.1 %	-0.20 [-0.37, -0.03]
3/61	2/20		8.9 %	-0.05 [-0.19, 0.09]
9/163	8/29		7.4 %	-0.22 [-0.39, -0.05]
683	354	•	80.8 %	-0.06 [-0.13, 0.01]
	bo)			
licable				
0/72	0/72	-	19.2 %	0.0 [-0.03, 0.03]
72	72	•		0.0 [-0.03, 0.03]
			1712 70	
	,			
(P = 1.0)				
755	426	•	100.0 %	-0.05 [-0.10, 0.01]
	,			
	0.00001); I ² =84%			
· ,				
$Chi^2 = 2.43, df = 1 (P =$	0.12), l ² =59%			
		-1 -0.5 0 0.5 1		
	(very potent) n/N 2/27 5/189 4/139 0/60 6/44 3/61 9/163 683 bid (very potent)), 35 (Place 26 (P = 0.097) 0 d (very potent)), 0 (Place bicable 0/72 72 d (very potent)), 0 (Place bicable 0/72 72 d (very potent)), 0 (Place 0/72 72 d (very potent)), 0 (Place 0/72 72 d (very potent)), 0 (Place 0/72 72 d (very potent)), 35 (Place 0/72 755 bid (very potent)), 35 (Place 0/72 756 0/72 757 0/72 757 0/72 0/72 758 0/72 756 0/72 756 0/72 756 0/72 756 0/72 757 0/72 756 0/72 757 0/72 756 0/72 756 0/72 757 0/72 756 0/72 756 0/72 757 0/72 756 0/72 757 0/72 757 0/72 757 0/72 756 0/72 756 0/72 757 0/72 756 0/72 756 0/72 757 0/72 756 0/72 757 0/72 757 0/72 757 0/72 757 0/72 757 0/72 757 0/72 757 0/72 757 0/72 757 0/72 757 0/72 757 0/72 7/72	(very potent) Placebo n/N n/N $2/27$ $2/27$ $5/189$ $4/33$ $4/139$ $4/140$ $0/60$ $0/60$ $6/44$ $15/45$ $3/61$ $2/20$ $9/163$ $8/29$ 683 354 bid (very potent)), 35 (Placebo) $2/20$ $Chi^2 = 33.33$, df = 6 (P<0.00001); l ² = 82% $66 (P = 0.097)$ 0 0 0 0 0 0 (very potent)), 0 (Placebo) blicable $0/72$ $0/72$ 72 $0 (P = 1.0)$ 755 426 243.07 , df = 7 (P<0.00001); l ² = 84% $66 (P = 0.12)$ $Chi^2 = 2.43$, df = 1 (P = 0.12), l ² = 59%	(very potent) Placebo Difference N/N n/N n/N $H_{Random,95\%}$ $2/27$ $2/27$ G $2/27$ $2/27$ G $2/27$ $2/27$ G $2/27$ $5/189$ $4/133$ \bullet $4/139$ $4/140$ \bullet \bullet $0/60$ $0/60$ \bullet \bullet $0/60$ $0/60$ \bullet \bullet $0/60$ $0/60$ \bullet \bullet $0/61$ $2/20$ \bullet \bullet $9/163$ $8/29$ \bullet \bullet $0/163$ $8/29$ \bullet \bullet $0/164$ $0/702$ \bullet \bullet $0/72$ $0/72$ $0/72$ \bullet $0/72$ $0/72$ 72 72 72 0	(very potent) Placebo Difference M- H,Random,95% Cl Weight M- H,Random,95% Cl $2/27$ $2/27$ 9.1% $2/27$ $2/27$ 9.1% $5/189$ $4/33$ 11.1% $4/139$ $4/140$ 18.3% $0/60$ $0/60$ 18.9% $6/44$ $15/45$ 7.1% $3/61$ $2/20$ 8.9% $9/163$ $8/29$ 7.4% 683 354 80.8% bid (very potent)), 35 (Placebo) $5h^2 = 33.3$, $df = 6$ (P<0.00001); P = 82%

Analysis 3.7. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Corticosteroid (very potent) n/N	Placebo n/N	Risk Difference M- H,Random,95% Cl	Weight	Risk Difference M,Random,95 Cl
I Clobetasol propionate	101 \$	1013			<u> </u>
Beutner 2006	0/27	0/27	+	1.7 %	0.0 [-0.07, 0.07]
Decroix 2004	1/189	0/33	+	4.4 %	0.01 [-0.04, 0.05]
Gottlieb 2003	0/139	1/140	-	20.7 %	-0.01 [-0.03, 0.01]
Jarratt 2006	0/60	0/60	+	7.8 %	0.0 [-0.03, 0.03]
Jorizzo 1997	1/44	1/45	+	2.1 %	0.00 [-0.06, 0.06]
Lebwohl 2002	0/61	0/20	<u> </u>	1.7 %	0.0 [-0.07, 0.07]
Lowe 2005	1/163	0/29		3.5 %	0.01 [-0.04, 0.05]
Subtotal (95% CI) Total events: 3 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.35	$f^2 = 0.56$, df = 6 (P = 1.0	<i>,</i>	ſ	41.9 %	0.00 [-0.02, 0.01]
2 Halcinonide Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable Test for overall effect: not appl		0 bo)			Not estimable
3 Halobetasol Bernhard 1991 (1)	0/100	0/100	+	21.3 %	0.0 [-0.02, 0.02]
Bernhard 1991(2)	0/72	0/72	+	11.1 %	0.0 [-0.03, 0.03]
Katz 1991b	0/110	0/110	-	25.7 %	0.0 [-0.02, 0.02]
Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch		,		58.1 %	0.0 [-0.01, 0.01]
Test for overall effect: Z = 0.0 Total (95% CI) Total events: 3 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.2; Test for subgroup differences:	965 (very potent)), 2 (Place ² = 0.59, df = 9 (P = 1.0 2 (P = 0.82)	00); l ² =0.0%	•	100.0 %	0.00 [-0.01, 0.01]

Favours corticosteroid (v potent) Favours placebo

Topical treatments for chronic plaque psoriasis (Review)

Analysis 3.8. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Corticosteroid (very potent)	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
I Clobetasol propionate					
Beutner 2006	0/27	0/27	+	3.2 %	0.0 [-0.07, 0.07]
Decroix 2004	0/189	1/33	+	3.2 %	-0.03 [-0.10, 0.04]
Gottlieb 2003	0/139	1/140	•	28.6 %	-0.01 [-0.03, 0.01]
Jarratt 2006	0/60	0/60	+	13.3 %	0.0 [-0.03, 0.03]
Jorizzo 1997	1/44	6/45		1.3 %	-0.11 [-0.22, 0.00]
Lebwohl 2002	0/61	0/20	+	3.2 %	0.0 [-0.07, 0.07]
Subtotal (95% CI)	520	325	4	52.9 %	-0.01 [-0.03, 0.01]
Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: not applicable Test for overall effect: not appl 3 Halobetasol Bernhard 1991 (1)		0 ebo) 0/100		29.1 %	Not estimable
Bernhard 1991(2)	0/72	0/72	-	17.9 %	0.0 [-0.03, 0.03]
Subtotal (95% CI) Total events: 0 (Corticosteroic Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: $Z = 0.0$	$i^2 = 0.0, df = 1 (P = 1.0)$,	·	47.1 %	0.0 [-0.02, 0.02]
Total (95% CI)	692	497		100.0 %	0.00 [-0.02, 0.01]
Total events: 1 (Corticosteroic Heterogeneity: Tau ² = 0.00; C Test for overall effect: $Z = 0.7$ (Test for subgroup differences:	$hi^2 = 8.09, df = 7 (P = 0)$ (P = 0.49)	$(0.32); ^2 = 3\%$			

Favours corticosteroid (v potent) Favours placebo

Topical treatments for chronic plaque psoriasis (Review)

Analysis 3.9. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

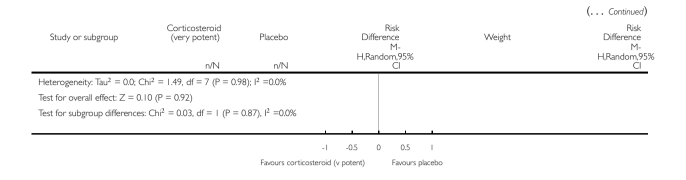
Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: 9 Adverse events (local)

Study or subgroup	Corticosteroid (very potent)	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,959 Cl
l Clobetasol propionate					
Beutner 2006	1/27	0/27		2.7 %	0.04 [-0.06, 0.13]
Decroix 2004	2/189	0/33	+	13.2 %	0.01 [-0.03, 0.05]
Gottlieb 2003	7/139	10/140	+	8.0 %	-0.02 [-0.08, 0.03]
Jarratt 2006	14/60	13/60		1.1 %	0.02 [-0.13, 0.17]
Jorizzo 1997	5/44	5/45	-	1.5 %	0.00 [-0.13, 0.13]
Lebwohl 2002	17/61	6/20		0.5 %	-0.02 [-0.25, 0.21]
Subtotal (95% CI)	520	325	•	27.0 %	0.00 [-0.03, 0.03]
2 Halcinonide Subtotal (95% CI) Total events: 0 (Corticostero Heterogeneity: not applicable Test for overall effect: not app 3 Halobetasol Bernhard 1991 (1)	e	0 bo) 0/100		67.0 %	Not estimable 0.0 [-0.02, 0.02]
Katz 1991b	7/110	7/110	+	6.0 %	0.0 [-0.06, 0.06]
Subtotal (95% CI) Total events: 7 (Corticostero Heterogeneity: Tau ² = 0.0; C		,		73.0 %	0.0 [-0.02, 0.02]
Test for overall effect: $Z = 0.0$, ,				
Total (95% CI)	730	535	+	100.0 %	0.00 [-0.02, 0.02]

(Continued . . .)

Topical treatments for chronic plaque psoriasis (Review)



Analysis 3.10. Comparison 3 Corticosteroid (very potent) versus placebo, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 3 Corticosteroid (very potent) versus placebo

Outcome: 10 Adverse events (systemic)

Study or subgroup	Corticosteroid (very potent)	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
l Clobetasol propionate					
Gottlieb 2003	0/139	1/140	•	23.4 %	-0.01 [-0.03, 0.01]
Jarratt 2006	0/60	0/60	+	8.8 %	0.0 [-0.03, 0.03]
Lebwohl 2002	17/61	6/20		0.2 %	-0.02 [-0.25, 0.21]
Subtotal (95% CI)	260	220	•	32.3 %	-0.01 [-0.02, 0.01]
Total events: 17 (Corticoster Heterogeneity: Tau ² = 0.0; C Test for overall effect: Z = 0. 2 Halcinonide Sudilovsky 1981	$Chi^2 = 0.21, df = 2 (P = 0)$,	-	14.7 %	0.0 [-0.02, 0.02]
Subtotal (95% CI)	78	78	•	14.7 %	0.0 [-0.02, 0.02]
Total events: 0 (Corticosterc Heterogeneity: not applicable Test for overall effect: $Z = 0$.	e	ebo)			
		Favours corticost	-0.5 -0.25 0 0.25 0.5 reroid (v potent) Favours placebo		(Continued)

(Continued ...)

Study or subgroup	Corticosteroid (very potent)	Placebo	Risk Difference M- H,Random,95%	Weight	(Continued) Risk Difference M- H,Random,95%
	n/N	n/N	Cl		Cl
3 Halobetasol					
Bernhard 1991 (1)	0/100	0/100	+	24.0 %	0.0 [-0.02, 0.02]
Katz 1991b	0/110	0/110	+	29.0 %	0.0 [-0.02, 0.02]
Subtotal (95% CI)	210	210	•	53.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Corticostero	oid (very potent)), 0 (Place	ebo)			
Heterogeneity: Tau ² = 0.0; C	$Chi^2 = 0.0, df = 1 (P = 1.0)$	0); I ² =0.0%			
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
Total (95% CI)	548	508	•	100.0 %	0.00 [-0.01, 0.01]
Total events: 17 (Corticoster	roid (very potent)), 7 (Pla	cebo)			
Heterogeneity: $Tau^2 = 0.0$; C	$2hi^2 = 0.5 I$, $df = 5 (P = 0.5)$	99); l ² =0.0%			
Test for overall effect: $Z = 0.2$	35 (P = 0.72)				
Test for subgroup differences	s: $Chi^2 = 0.26$, $df = 2$ (P =	= 0.88), I ² =0.0%			

-0.5 -0.25 0 0.25 0.5

Favours corticosteroid (v potent) Favours placebo

Analysis 4.2. Comparison 4 Dithranol versus placebo, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 4 Dithranol versus placebo

Outcome: 2 TSS

Study or subgroup	Dithranol N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
Buckley 1978	8	-0.54 (0.29)	8	-0.26 (0.14)	-#-	22.7 %	-1.16 [-2.25, -0.08]
Grattan 1997 (P)	12	1.2 (1.77)	12	4.1 (1.59)	+	27.3 %	-1.66 [-2.62, -0.71]
Jekler 1992	27	0.99 (0.47)	27	1.3 (0.42)	-	50.0 %	-0.69 [-1.24, -0.14]
Total (95% CI) Heterogeneity: Tau ² = Test for overall effect:			47 0.20); I ² =37%		•	100.0 %	-1.06 [-1.66, -0.46]
Test for subgroup diffe	`	,					
				F	-10 -5 0 5 avours dithranol Favours plac	10 cebo	

Topical treatments for chronic plaque psoriasis (Review)

Analysis 4.5. Comparison 4 Dithranol versus placebo, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 4 Dithranol versus placebo

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Dithranol		Placebo		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% C	1	IV,Random,95% CI
Buckley 1978	8	-0.54 (0.29)	8	-0.26 (0.14)	=	22.7 %	-1.16 [-2.25, -0.08]
Grattan 1997 (P)	12	1.2 (1.77)	12	4.1 (1.59)	-	27.3 %	-1.66 [-2.62, -0.71]
Jekler 1992	27	0.99 (0.47)	27	1.3 (0.42)	•	50.0 %	-0.69 [-1.24, -0.14]
Total (95% CI) Heterogeneity: Tau ² =	47 = 0.11; Chi ² = 3	8.19, df = 2 (P = 0.2	47 0); I ² =37%	6	•	100.0 %	-1.06 [-1.66, -0.46]
Test for overall effect:	Z = 3.45 (P =	0.00056)					
Test for subgroup diffe	erences: Not ap	plicable					
					<u> </u>		
					-20 -10 0 10	20	

Favours dithranol Favours placebo

Analysis 4.6. Comparison 4 Dithranol versus placebo, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 4 Dithranol versus placebo

Outcome: 6 Total withdrawals

Study or subgroup	Dithranol	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
Buckley 1978	2/10	2/10		7.3 %	0.0 [-0.35, 0.35]
Grattan 1997 (P)	0/12	0/12	+	40.9 %	0.0 [-0.15, 0.15]
Jekler 1992	3/30	3/30	+	38.8 %	0.0 [-0.15, 0.15]
Volden 1992	1/10	1/10	- _	12.9 %	0.0 [-0.26, 0.26]
Total (95% CI)	62	62	+	100.0 %	0.0 [-0.09, 0.09]
Total events: 6 (Dithranol	I), 6 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$	0; Chi ² = 0.0, df = 3 (P	= 1.00); 1 ² =0.0%			
Test for overall effect: Z =	= 0.0 (P = 1.0)				
Test for subgroup differer	nces: Not applicable				
			-I -0.5 0 0.5 I		

- I -0.5 0 0.5 I Favours dithranol Favours placebo

Analysis 4.7. Comparison 4 Dithranol versus placebo, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 4 Dithranol versus placebo

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Dithranol	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
Grattan 1997 (P)	0/12	0/12		13.7 %	0.0 [-0.15, 0.15]
Jekler 1992	0/30	0/30	-	76.4 %	0.0 [-0.06, 0.06]
Volden 1992	0/10	0/10	-	9.9 %	0.0 [-0.17, 0.17]
Total (95% CI)	52	52	•	100.0 %	0.0 [-0.05, 0.05]
Total events: 0 (Dithrano	I), 0 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$	0; Chi ² = 0.0, df = 2 (P	= 1.00); l ² =0.0%			
Test for overall effect: Z =	= 0.0 (P = 1.0)				
Test for subgroup differer	nces: Not applicable				
			-I -0.5 0 0.5 I		

Favours dithranol Favours placebo

Analysis 4.8. Comparison 4 Dithranol versus placebo, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 4 Dithranol versus placebo

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Dithranol	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
Grattan 1997 (P)	0/12	0/12	-	58.1 %	0.0 [-0.15, 0.15]
Volden 1992	0/10	0/10	+	41.9 %	0.0 [-0.17, 0.17]
Total (95% CI)	22	22	+	100.0 %	0.0 [-0.11, 0.11]
Total events: 0 (Dithranol), 0 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$); $Chi^2 = 0.0$, $df = 1$ (P	= 1.00); 1 ² =0.0%			
Test for overall effect: Z =	= 0.0 (P = 1.0)				
Tast for subgroup differen	ices: Not applicable				

- I -0.5 0 0.5 I Favours dithranol Favours placebo

Analysis 4.9. Comparison 4 Dithranol versus placebo, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 4 Dithranol versus placebo

Outcome: 9 Adverse events (local)

Study or subgroup	Dithranol	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl_
Buckley 1978	5/7	2/7		29.5 %	0.43 [-0.04, 0.90]
Jekler 1992	0/30	0/30	•	37.1 %	0.0 [-0.06, 0.06]
Volden 1992	4/10	0/10		33.4 %	0.40 [0.08, 0.72]
Total (95% CI)	47	47	-	100.0 %	0.26 [-0.30, 0.82]
Total events: 9 (Dithrano	I), 2 (Placebo)				
Heterogeneity: $Tau^2 = 0.2$	22; Chi ² = 25.44, df = 2	2 (P<0.00001); I ² =92%	,)		
Test for overall effect: Z =	= 0.91 (P = 0.36)				
Test for subgroup differer	nces: Not applicable				
			-2 -1 0 1 2		
		F	avours dithranol Favours placebo		

Analysis 4.10. Comparison 4 Dithranol versus placebo, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 4 Dithranol versus placebo

Outcome: 10 Adverse events (systemic)

Study or subgroup	Dithranol	Placebo	Risk Difference M- H.Random,95%	Weight	Risk Difference M- H.Random,95%
	n/N	n/N	Cl		Cl
Buckley 1978	2/10	2/10	-	100.0 %	0.0 [-0.35, 0.35]
Total (95% CI)	10	10	+	100.0 %	0.0 [-0.35, 0.35]
Total events: 2 (Dithranol), 2 (Placebo)				
Heterogeneity: not applic	able				
Test for overall effect: Z =	= 0.0 (P = 1.0)				
Test for subgroup differen	ices: Not applicable				
			-2 -I 0 I 2		
			Favours dithranol Favours placebo		

Analysis 5.1. Comparison 5 Vitamin D combination products versus placebo, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: I IAGI

Sto Mea Differenc	Weight	Std. Mean Difference		Placebo		Vitamin D combina- tion	Study or subgroup
IV,Random,95% (IV,Random,95% Cl	Mean(SD)	Ν	Mean(SD)	Ν	
				ce daily	dipropionate, on	etamethasone	I Combination calcipotriol/b
-0.79 [-1.20, -0.38	14.5 %	-	2.64 (0.62)	28	1.85 (1.05)	150	Fleming 2010 (P)
-1.58 [-1.82, -1.33	17.2 %	-	-1.88 (1.14)	206	-3.59 (1)	150	Guenther 2002 (P)
-1.31 [-1.51, -1.12	17.8 %	-	2.68 (0.89)	157	1.5 (0.9)	490	Kaufmann 2002 (P)
-1.02 [-1.32, -0.72	16.3 %	-	2.75 (0.78)	64	1.84 (0.93)	171	Langley 2011 (P)
-1.21 [-1.50, -0.91	65. 7 %	•		455		961	Subtotal (95% CI)
); l ² =79%	df = 3 (P = 0.003)	$Chi^2 = 14.26,$	Heterogeneity: Tau ² = 0.07;
					01)	96 (P < 0.000	Test for overall effect: $Z = 7.9$
				ce daily	dipropionate, tw	etamethasone	2 Combination calcipotriol/b
-1.81 [-2.03, -1.59	17.4 %	-	-1.88 (1.14)	206	-3.79 (0.97)	234	Guenther 2002 (P)
-2.01 [-2.27, -1.75	16.9 %	-	-1.92 (1.07)	107	-3.84 (0.91)	301	Papp 2003 (P)
-1.90 [-2.09, -1.71	34.3 %	•		313		535	Subtotal (95% CI)
				² =22%	f = I (P = 0.26);	Chi ² = 1.27, d	Heterogeneity: $Tau^2 = 0.00;$
					001)	9.38 (P < 0.00	Test for overall effect: Z = 19
-1.44 [-1.76, -1.12	100.0 %	•		768		1496	Total (95% CI)
			,	I); I ² =89%	df = 5 (P<0.0000	Chi ² = 47.32,	Heterogeneity: $Tau^2 = 0.14;$
					01)	73 (P < 0.000	Test for overall effect: $Z = 8.7$
), I ² =93%	7, df = 1 (P = 0.00	:: Chi ² = 14.67	Test for subgroup differences

Favours vitamin D combination

Favours placebo

Analysis 5.3. Comparison 5 Vitamin D combination products versus placebo, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: 3 PASI

	Vitamin D combina- tion N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% Cl
I Combination calcipotriol/bet	tamethasone	e dipropionate, or	ice daily				
Fleming 2010 (P)	147	3.1 (2.8)	28	5.5 (3.5)	-	13.9 %	-0.82 [-1.23, -0.40]
Guenther 2002 (P)	150	3 (2.5)	207	7.7 (5.5)	-	17.4 %	-1.04 [-1.27, -0.82]
Kaufmann 2002 (P)	490	-0.71 (0.26)	157	-0.23 (0.34)	•	17.8 %	-1.70 [-1.91, -1.50]
Langley 2011 (P)	171	3.58 (2.98)	64	6.47 (3.56)	-	16.1 %	-0.92 [-1.21, -0.62]
Subtotal (95% CI)	958		456		•	65.2 %	-1.14 [-1.57, -0.70]
Heterogeneity: $Tau^2 = 0.17$; C	hi ² = 31.51,	df = 3 (P<0.0000); ² =909	%			
Test for overall effect: $Z = 5.13$	3 (P < 0.000	01)					
2 Combination calcipotriol/bet	tamethasone	dipropionate, tw	rice daily				
Guenther 2002 (P)	234	2.7 (2.5)	207	7.7 (5.5)	-	17.8 %	-1.19 [-1.40, -0.99]
Papp 2003 (P)	301	-0.73 (0.25)	107	-0.29 (0.31)	-	17.0 %	-1.65 [-1.89, -1.40]
Subtotal (95% CI)	535		314		•	34.8 %	-1.41 [-1.86, -0.97]
Heterogeneity: $Tau^2 = 0.09$; C	$hi^2 = 7.65, c$	f = (P = 0.01);	l ² =87%				
Test for overall effect: $Z = 6.25$	5 (P < 0.000	01)					
Total (95% CI)	1493	,	77 0		•	100.0 %	-1.24 [-1.53, -0.95]
Heterogeneity: $Tau^2 = 0.11$; C	hi ² = 40.35,	df = 5 (P<0.0000); l ² =889	%			
Test for overall effect: $Z = 8.39$	9 (P < 0.000	01)					
Test for subgroup differences:	$Chi^2 = 0.76,$	df = 1 (P = 0.38)), l ² =0.0%				

-10 -5 0 5 10

Favours vitamin D combination Favours placebo

Analysis 5.4. Comparison 5 Vitamin D combination products versus placebo, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: 4 PAGI

Study or subgroup	Vitamin D combina- tion		Placebo			C	St Mea Differend	an		Std. Mean Difference
	Ν	Mean(SD) N		Mean(SD)	IV,Random,95% CI				IV,Random,95% CI	
I Combination calcipotr	iol/betamethasone d	ipropionate, once daily								
Langley 2011 (P)	171	1.82 (0.94)	64	2.48 (0.99)			+			-0.69 [-0.98, -0.40]
2 Combination calcipotr	iol/betamethasone d	propionate, twice daily	/					1		
					-10	-5	0	5	10	
				Favours vitar	nin D comb	pination	Fa	ivours	placebo	

Analysis 5.5. Comparison 5 Vitamin D combination products versus placebo, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D combina- tion		Placebo		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
I Combination calcipotriol/b	etamethasone	e dipropionate, or	nce daily				
Fleming 2010 (P)	150	1.85 (1.05)	28	2.64 (0.62)		14.5 %	-0.79 [-1.20, -0.38]
Guenther 2002 (P)	150	-3.59 (1)	206	-1.88 (1.14)	•	17.2 %	-1.58 [-1.82, -1.33]
Kaufmann 2002 (P)	490	1.5 (0.9)	157	2.68 (0.89)	-	17.8 %	-1.31 [-1.51, -1.12]
Langley 2011 (P)	171	1.84 (0.93)	64	2.75 (0.78)	+	16.3 %	-1.02 [-1.32, -0.72]
Subtotal (95% CI)	961		455		•	65.7 %	-1.21 [-1.50, -0.91]
Heterogeneity: $Tau^2 = 0.07$;	Chi ² = 14.26,	df = 3 (P = 0.00	3); I ² =79%				
Test for overall effect: $Z = 7$.	96 (P < 0.000	01)	,				
2 Combination calcipotriol/b	etamethasone	e dipropionate, tw	vice daily				
Guenther 2002 (P)	234	-3.79 (0.97)	206	-1.88 (1.14)	-	17.4 %	-1.81 [-2.03, -1.59]
Papp 2003 (P)	301	-3.84 (0.91)	107	-1.92 (1.07)	•	16.9 %	-2.01 [-2.27, -1.75]
Subtotal (95% CI)	535		313		•	34.3 %	-1.90 [-2.09, -1.71]
Heterogeneity: $Tau^2 = 0.00;$	$Chi^2 = 1.27, c$	f = 1 (P = 0.26);	l ² =22%				
Test for overall effect: $Z = 19$	9.38 (P < 0.00	001)					
Total (95% CI)	1496		768		•	100.0 %	-1.44 [-1.76, -1.12]
Heterogeneity: Tau ² = 0.14;	Chi ² = 47.32,	df = 5 (P<0.000	01); I ² =899	%			
Test for overall effect: $Z = 8$.	73 (P < 0.000	01)					
Test for subgroup differences	s: $Chi^2 = 14.6$	7, df = 1 (P = 0.0	0), I ² =93%				
						1	
				-4	-2 0 2	4	
				Favours vitamin D	combination Eavours pla	icebo	

Favours vitamin D combination Favours placebo

Analysis 5.6. Comparison 5 Vitamin D combination products versus placebo, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D combina- tion	Placebo	Risk Difference M-	Weight	Risk Difference H,Random,95% Cl	
	n/N	n/N	H,Random,95% Cl			
I Combination calcipotriol/beta	amethasone dipropio	nate, once daily				
Fleming 2010 (P)	13/162	12/40		7.8 %	-0.22 [-0.37, -0.07]	
Guenther 2002 (P)	11/152	34/208	-	19.3 %	-0.09 [-0.16, -0.03]	
Kaufmann 2002 (P)	13/490	25/157	-	20.5 %	-0.13 [-0.19, -0.07]	
Langley 2011 (P)	12/183	27/91	-	12.9 %	-0.23 [-0.33, -0.13]	
Subtotal (95% CI)	987	496	•	60.5 %	-0.15 [-0.22, -0.09]	
Test for overall effect: Z = 4.69 2 Combination calcipotriol/bet Guenther 2002 (P)	()	nate, twice daily 34/208		20.1 %	-0.08 [-0.14, -0.02]	
Papp 2003 (P)	16/304	12/108		19.4 %	-0.06 [-0.12, 0.01]	
Subtotal (95% CI) Total events: 35 (Vitamin D cor Heterogeneity: Tau ² = 0.0; Chi	$^{2} = 0.30, df = 1 (P =$,	•	39.5 %	-0.07 [-0.12, -0.03]	
Test for overall effect: Z = 3.17 Total (95% CI)	(P = 0.0015) 1528	812	•	100.0 %	-0.12 [-0.17, -0.07]	
Total events: 84 (Vitamin D con Heterogeneity: Tau ² = 0.00; Ch Test for overall effect: $Z = 4.89$	mbination), 144 (Place ni ² = 12.27, df = 5 (P	ebo)		100.0 /0	-0.12 [-0.17, -0.07]	

Favours vitamin D combination Favours placebo

Analysis 5.7. Comparison 5 Vitamin D combination products versus placebo, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D combina- tion	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
I Combination calcipotriol/beta	methasone dipropior	nate, once daily			
Guenther 2002 (P)	0/151	21/208	-	25.4 %	-0.10 [-0.14, -0.06]
Kaufmann 2002 (P)	3/490	12/157	-	25.5 %	-0.07 [-0.11, -0.03]
Langley 2011 (P)	3/183	4/91	-	23.5 %	-0.03 [-0.07, 0.02]
Subtotal (95% CI)	824	456	•	74.4 %	-0.07 [-0.11, -0.03]
Total events: 6 (Vitamin D com	oination), 37 (Placebo	o)			
Heterogeneity: $Tau^2 = 0.00$; Ch	i ² = 5.36, df = 2 (P =	= 0.07); l ² =63%			
Test for overall effect: $Z = 3.21$	(P = 0.00 3)				
2 Combination calcipotriol/beta	methasone dipropio	nate, twice daily			
Guenther 2002 (P)	1/235	21/208	-	25.6 %	-0.10 [-0.14, -0.05]
Subtotal (95% CI)	235	208	•	25.6 %	-0.10 [-0.14, -0.05]
Total events: I (Vitamin D com	pination), 21 (Placebo	o)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 4.54$	(P < 0.00001)				
Total (95% CI)	1059	664	•	100.0 %	-0.07 [-0.11, -0.04]
Total events: 7 (Vitamin D com	pination), 58 (Placebo	o)			
Heterogeneity: $Tau^2 = 0.00$; Ch	i ² = 6.67, df = 3 (P =	= 0.08); l ² =55%			
Test for overall effect: $Z = 4.58$	(P < 0.00001)				
Test for subgroup differences: C	$hi^2 = 0.98, df = 1 (P$	= 0.32), I ² =0.0%			
			-I -0.5 0 0.5 I		
		Favours vitamir	n D combination Favours placebo	1	

Analysis 5.8. Comparison 5 Vitamin D combination products versus placebo, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: 8 Withdrawals due to treatment failure

tion	Placebo	Risk Difference	Weight	Risk Difference	
n/N	n/N	M- H,Random,95% Cl		M- H,Random,95% Cl	
thasone dipropio	nate, once daily				
0/151	19/208	•	49.5 %	-0.09 [-0.13, -0.05]	
151	208	•	49.5 %	-0.09 [-0.13, -0.05]	
< 0.00001)	,		50.5 %	-0.09 [-0.13, -0.05]	
235	208	•	50.5 %	-0.09 [-0.13, -0.05]	
<i>,</i> , ,)				
386	416	•	100.0 %	-0.09 [-0.12, -0.06]	
ition), 38 (Placebo	o)				
0.02, df = 1 (P =	0.88); I ² =0.0%				
< 0.00001)					
= 0.02, df = 1 (P	= 0.88), I ² =0.0%				
	thasone dipropion 0/151 151 (tion), 19 (Placebo < 0.00001) thasone dipropion 1/235 235 (tion), 19 (Placebo 386 (tion), 38 (Placebo 0.02, df = 1 (P = < 0.00001)	thasone dipropionate, once daily 0/151 19/208 151 208 Ition), 19 (Placebo) < 0.00001) thasone dipropionate, twice daily 1/235 19/208 235 208 Ition), 19 (Placebo) = 0.000020) 386 416 Ition), 38 (Placebo) 0.02, df = 1 (P = 0.88); 1 ² = 0.0%	n/N Cl thasone dipropionate, once daily $0/151$ 19/208 151 208 • ition), 19 (Placebo) • • < 0.00001)	H,Random,95% Cl n/N n/N Cl thasone dipropionate, once daily 0/151 19/208 49.5 % 151 208 49.5 % tion), 19 (Placebo) 50.5 % 50.5 % 235 208 50.5 % tion), 19 (Placebo) 50.5 % 100.0 % attion), 19 (Placebo) 100.0 % attion), 19 (Placebo) 100.0 % attion), 38 (Placebo) 0.02, df = 1 (P = 0.88); 1 ² = 0.0% output 100.0 %	

Favours vitamin D combination Favours placebo

Analysis 5.9. Comparison 5 Vitamin D combination products versus placebo, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D combina- tion	Placebo		Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N		H,Random,95% Cl		H,Random,95% Cl
I Combination calcipotriol/beta	amethasone dipropio	nate, once daily				
Fleming 2010 (P)	12/160	10/40		_ 	4.8 %	-0.18 [-0.32, -0.03]
Guenther 2002 (P)	15/151	26/208		•	19.7 %	-0.03 [-0.09, 0.04]
Kaufmann 2002 (P)	29/490	21/157		-	25.0 %	-0.07 [-0.13, -0.02]
Langley 2011 (P)	16/182	4/9		-	12.4 %	-0.07 [-0.15, 0.02]
Subtotal (95% CI)	983	496		•	61.9 %	-0.07 [-0.11, -0.02]
Heterogeneity: $Tau^2 = 0.00$; Ch Test for overall effect: $Z = 2.95$ 2 Combination calcipotriol/beta	(P = 0.0032)	,				
2 Combination calcipotriol/beta Guenther 2002 (P)	amethasone dipropio 25/235	nate, twice daily 26/208		+	23.2 %	-0.02 [-0.08, 0.04]
Papp 2003 (P)	30/304	17/108			15.0 %	-0.06 [-0.14, 0.02]
Subtotal (95% CI)	539	316		•	38.1 %	-0.03 [-0.08, 0.01]
Total events: 55 (Vitamin D cor Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: $Z = 1.41$	mbination), 43 (Placeb 2 = 0.66, df = 1 (P =	00)				
Total (95% CI)	1522	812		•	100.0 %	-0.05 [-0.08, -0.02]
Total events: 127 (Vitamin D cc Heterogeneity: Tau ² = 0.00; Ch Test for overall effect: $Z = 3.36$ Test for subgroup differences: C	$hi^2 = 5.56, df = 5 (P = 0.00077)$	= 0.35); I ² = I 0%				
			-1 -0.5	0 0.5 1		
		Favours vitami	in D combinati		bo	

Analysis 5.10. Comparison 5 Vitamin D combination products versus placebo, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 5 Vitamin D combination products versus placebo

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D combina- tion	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
I Combination calcipotriol/beta	methasone dipropion	ate, once daily			
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D comb	pination), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applic	able				
2 Combination calcipotriol/beta	methasone dipropion	ate, twice daily			
Papp 2003 (P)	0/304	0/108		100.0 %	0.0 [-0.0 , 0.0]
Subtotal (95% CI)	304	108		100.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Vitamin D comb	pination), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ (F	P = 1.0)				
Total (95% CI)	304	108		100.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Vitamin D comb	pination), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ (F	P = 1.0				
Test for subgroup differences: N	ot applicable				
			-I -0.5 0 0.5 I		
		Favours vitamin	D combination Favours placebo	c	

Favours vitamin D combination Favours placebo

Analysis 6.1. Comparison 6 Other treatment versus placebo, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: I IAGI

Study or subgroup C	Other treatment		Placebo		Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% Cl	IV,Random,95% CI
I Aloe vera extract 0.5% hy	drophilic cream, thr	ee times per day				
2 Anti-IL-8 monoclonal antib	body cream					
Jin 2001	45	-1.27 (1.01)	44	-0.73 (0.79)	+	-0.59 [-1.01, -0.16]
3 Betamethasone 17-valerat	e 21-acetate plus tr	etinoin plus salicylic ac	id			
Santoianni 2001	42	-2.36 (0.69)	39	-1.79 (0.8)	+	-0.76 [-1.21, -0.31]
4 Caffeine (topical) 10%, TD)					
5 Calcipotriene 0.005% oint	ment + nicotinamic	le 0.05% or 0.1% or 0.1	7% or 1.4%, t	wice daily		
6 Dead Sea salts emollient le	otion					
7 Fish oil plus occlusion						
8 Herbal skin care (Dr Mich	aels cleansing gel,	ointment and skin con	iditioner), twi	ce daily		
9 Hexafluoro-1,25-dihydrox	yvitamin D3					
Durakovic 2001	15	-3.3 (0.71)	15	-2.8 (0.86)	-	-0.62 [-1.35, 0.12]
10 Indigo naturalis 1.4% oint	tment					
Lin 2008	34	-4.18 (1.22)	34	-1.68 (1.09)	+	-2.14 [-2.74, -1.53]
I I Kukui nut oil, TD						
Brown 2005	13	1.7 (0.9)	П	1.7 (0.8)	+	0.0 [-0.80, 0.80
12 Mahonia aquifolium	ı⁄ (Reli⊃va™), twice	daily				
13 Methotrexate gel						
Sutton 2001	39	-2.34 (1.2)	41	-1.78 (0.72)	+	-0.56 [-1.01, -0.12]
14 Mycophenolic acid ointm	nent					
15 NG-monomethyl-L-argin	iine (L-NMMA) crea	ım				
16 Nicotinamide 1.4%, twice	e daily					
17 Oleum horwathiensis (Ps	soricur)					
Lassus 1991	19	-2.21 (1.69)	23	-2.17 (1.72)	+	-0.02 [-0.63, 0.58]
18 Omega-3-polyunsaturate	ed fatty acids ointme	ent				
19 Platelet aggregation activ	ating factor (PAF)(R	o 24-0238)				
Wolska 1995	40	-3.13 (0.79)	40	-3.08 (0.73)	+	-0.07 [-0.50, 0.37]
		. /		. /		
					-10 -5 0 5	10
				Favou	rs other treatment Favours pl	acebo
						(Continued

Study or subgroup	Other treatment		Placebo	Std. Mean Difference				(Continued) Std. Mean Difference		
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% Cl				IV,Random,95% CI	
20 Polymyxin B cream	200,000 U/g									
21 PTH (1-34) in Nov	asome A liposomal crea	m, twice daily								
22 Sirolimus (topical),	2.2% for 6 wks, then 8% f	or a further 6 wks								
23 Tacrolimus ointmer	nt									
24 Tar										
25 Tazarotene										
26 Theophylline 1% oi	intment, twice daily									
					-10	-5 0	5	10		
				Favour	s other treat	ment	Favours	placebo		

Analysis 6.2. Comparison 6 Other treatment versus placebo, Outcome 2 TSS.

<u>.</u>						Std. Mean		Std. Mean
Study or subgroup	Other treatment N	Mean(SD)	Placebo N	Mean(SD)	_	ifference dom,95% Cl	Weight	Difference IV,Random,95% CI
I Aloe vera extract 0.5%		· /		cu(3D)	. •,1 \			
Subtotal (95% CI)		ee umes per day	0					Not estimable
Heterogeneity: not appli	cable							
Test for overall effect: no	t applicable							
2 Anti-IL-8 monoclonal a	antibody cream							
Jin 2001	45	4.78 (3.2)	44	7.09 (3.31)		-	100.0 %	-0.70 [-1.13, -0.27]
Subtotal (95% CI)	45		44			•	100.0 %	-0.70 [-1.13, -0.27]
Heterogeneity: not appli	cable							
Test for overall effect: Z	= 3.22 (P = 0.0013)							
3 Betamethasone 17-vale	erate 21-acetate plus tr	retinoin plus salicy	/lic acid					
				1				
				-10	-5	0 5 I	0	
				Favours other	treatment	Favours place	ebo	
								(Continued

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Review: Topical treatments for chronic plaque psoriasis Comparison: 6 Other treatment versus placebo

					Std. Mean		(Continu St Mea
Study or subgroup	Other treatment N	Mean(SD)	Placebo N	Mean(SD)	Difference IV,Random,95% Cl	Weight	Differenc IV,Random,95% (
Subtotal (95% CI)	0	()	0	()			Not estimabl
Heterogeneity: not applica			-				
Test for overall effect: not							
4 Caffeine (topical) 10%, ⁻							
Subtotal (95% CI)	0		0				Not estimab
Heterogeneity: not applica	able						
Test for overall effect: not	applicable						
5 Calcipotriene 0.005% oi	intment + nicotinamie	de 0.05% or 0.	% or 0.7% (or 1.4%, twice daily	,		
Levine 2010 (P)	144	3.15 (1.94)	48	4.08 (1.93)	-	100.0 %	-0.48 [-0.81, -0.15
Subtotal (95% CI)	144		48		•	100.0 %	-0.48 [-0.81, -0.15
Heterogeneity: not applica	able						
Test for overall effect: Z =	: 2.84 (P = 0.0045)						
6 Dead Sea salts emollien	t lotion						
Subtotal (95% CI)	0		0				Not estimab
Heterogeneity: not applica	able						
Test for overall effect: not	applicable						
7 Fish oil plus occlusion							
Escobar 1992	25	1.74 (1.94)	25	3.81 (1.94)	+	100.0 %	-1.05 [-1.64, -0.46
Subtotal (95% CI)	25		25		•	100.0 %	-1.05 [-1.64, -0.46
Heterogeneity: not applica	able						
Test for overall effect: Z =	: 3.46 (P = 0.00053)						
8 Herbal skin care (Dr Mi	chaels cleansing gel,	ointment and	skin conditio	oner), twice daily			
Subtotal (95% CI)	0		0				Not estimab
Heterogeneity: not applica	able						
Test for overall effect: not	applicable						
9 Hexafluoro- I ,25-dihydn	oxyvitamin D3, twice	daily					
Durakovic 2001	15	2.4 (4.9)	15	8.1 (4.9)		100.0 %	-1.13 [-1.91, -0.3
Subtotal (95% CI)	15		15		•	100.0 %	-1.13 [-1.91, -0.35
Heterogeneity: not applica	able						
Test for overall effect: Z =							
10 Indigo naturalis 1.4% o	interant						
Lin 2007	I0	3.7 (2)	10	7.2 (1.4)		19.7 %	-1.94 [-3.05, -0.84
		. ,		. ,	_		
Lin 2008	34	6.3 (4.28)	34	12.8 (3.92)	-	80.3 %	-1.57 [-2.11, -1.02
Subtotal (95% CI)	44		44		•	100.0 %	-1.64 [-2.13, -1.15
Heterogeneity: $Tau^2 = 0.0$); $Chi^2 = 0.36$, $df = 1$	$(P = 0.55); ^2 =$	=0.0%				
	= 6.56 (P < 0.00001)						
lest for overall effect: \angle =							
lest for overall effect: Z = I I Kukui nut oil, TD		5.55 (2.26)	11	4.77 (2.26)		100.0 %	0.33 [-0.48, 1.14
	13	0.00 (2.20)				100.0.0/	
I I Kukui nut oil, TD Brown 2005		5155 (1126)	11		*		033 .048 . 14
I I Kukui nut oil, TD Brown 2005 Subtotal (95% CI)	13	5155 (2125)	11		•	100.0 %	0.33 [-0.48, 1.14
I I Kukui nut oil, TD Brown 2005	13	0.00 (2.20)	11		•	100.0 %	0.33 [-0.48, 1.14
I I Kukui nut oil, TD Brown 2005 Subtotal (95% CI)	13		11	-10	-5 0 5	100.0 %	0.33 [-0.48, 1.14

Study or subgroup Other t	reatment N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	(Continued Std. Mean Difference IV,Random,95% CI
Test for overall effect: Z = 0.81 (P =	= 0.42)						
12 <i>Mahonia aquifolium</i> (Reli Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not applicable	0	e daily	0				Not estimable
3 Methotrexate gel Sutton 2001	41	-2.98 (3.93)	41	-1.32 (2.77)	-	100.0 %	-0.48 [-0.92, -0.04]
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 2.16 (P =	41 = 0.031)		41		•	100.0 %	-0.48 [-0.92, -0.04]
14 Mycophenolic acid ointment Geilen 2000	7	5.13 (0.73)	7	6.32 (0.81)		100.0 %	-1.44 [-2.67, -0.22
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 2.32 (P =	7 = 0.021)		7		•	100.0 %	-1.44 [-2.67, -0.22]
I 5 NG-monomethyl-L-arginine (L-N Ormerod 2000	NMMA) cre 17	eam 9.12 (3.55)	17	8.82 (4.22)	-	100.0 %	0.08 [-0.60, 0.75
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 0.22 (P =	17 = 0.83)		17		+	100.0 %	0.08 [-0.60, 0.75
16 Nicotinamide 1.4%, twice daily Levine 2010 (P)	48	3.69 (1.93)	48	4.08 (1.93)		100.0 %	-0.20 [-0.60, 0.20
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 0.98 (P =	48	5.67 (175)	48	100 (175)	Ī	100.0 %	-0.20 [-0.60, 0.20]
17 Oleum horwathiensis (Psoricur Lassus 1991)	-0.74 (0.23)	23	-0.56 (0.23)	-	100.0 %	-0.77 [-1.40, -0.14
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 2.38 (P =	19 = 0.017)		23		•	100.0 %	-0.77 [-1.40, -0.14
18 Omega-3-polyunsaturated fatty a Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not applicable	0	lent	0				Not estimable
19 Platelet aggregation activating fac Subtotal (95% CI) Heterogeneity: not applicable	tor (PAF)(I 0	Ro 24-0238)	0				Not estimable

(... Continued)

Study or subgroup O	ther treatment		Placebo		Std. Mean Difference	Weight	Sto Mea Differenc
stady of subgroup	N	Mean(SD)	N	Mean(SD)	IV,Random,95% CI	* Volgite	IV,Random,95% C
Test for overall effect: not app	licable						
20 Polymyxin B cream 200,00	0 U/g						
Stutz 1996	15	2.5 (1.5)	15	2.3 (1.5)		100.0 %	0.13 [-0.59, 0.85
Subtotal (95% CI)	15		15		+	100.0 %	0.13 [-0.59, 0.85
Heterogeneity: not applicable Test for overall effect: Z = 0.3	5 (P = 0.72)						
21 PTH (1-34) in Novasome .	A liposomal cre	eam, twice daily					
Holick 2003	15	-0.67 (0.23)	15	-0.18 (0.18)		100.0 %	-2.31 [-3.26, -1.36
Subtotal (95% CI)	15		15		•	100.0 %	-2.31 [-3.26, -1.36
Heterogeneity: not applicable Test for overall effect: Z = 4.7	6 (P < 0.00001)						
22 Sirolimus (topical), 2.2% for Ormerod 2005	r 6 wks, then 8% 22	for a further 6 w 9.1 (4.8)	/ks 22	11.2 (5.8)	-	100.0 %	-0.39 [-0.98, 0.21
		7.1 (4.0)		11.2 (3.0)			-
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 1.2	22 7 (P = 0.20)		22			100.0 %	-0.39 [-0.98, 0.21
23 Tacrolimus ointment							
Zonneveld 1998 (P)	24	4.7 (1.73)	23	4.6 (1.73)	-	100.0 %	0.06 [-0.52, 0.63
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: $Z = 0.1^{\circ}$	24 9 (P = 0.85)		23		•	100.0 %	0.06 [-0.52, 0.63
24 Tar							
Kanzler 1993	18	-3.25 (2.03)	18	-2.36 (1.86)	H	100.0 %	-0.45 [-1.11, 0.22
Subtotal (95% CI)	18		18		•	100.0 %	-0.45 [-1.11, 0.22
Heterogeneity: not applicable Test for overall effect: Z = 1.3	2 (P = 0.19)						
25 Tazarotene Weinstein 1996	211	3.8 (1.73)	107	5.3 (1.73)		100.0 %	-0.86 [-1.11, -0.62
Subtotal (95% CI)	211	5.0 (1.75)	107	5.5 (1.75)	•		-0.86 [-1.11, -0.62
Heterogeneity: not applicable Test for overall effect: Z = 7.0			10/			100.0 %	-0.00 [-1.11, -0.02
26 Theophylline 1% ointment,	twice daily						
Subtotal (95% CI) Heterogeneity: not applicable	0		0				Not estimabl
Test for overall effect: not app Test for subgroup differences:		= 16 (P = 0.00),	l ² =75%				
						ı	
Test for subgroup differences:	Chi ² = 63.44, df	= 16 (P = 0.00),	² =75%	-10 Favours other		• 10 cebo	

Analysis 6.3. Comparison 6 Other treatment versus placebo, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: 3 PASI

Study or subgroup	Other treatment		Placebo		Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% C	IV,Random,95% CI
Aloe vera extract 0.5%	nydrophilic cream, three tir	mes per day				
Syed 1996	30	2.2 (3.76)	30	8.2 (3.76)	+	-1.58 [-2.16, -0.99]
2 Anti-IL-8 monoclonal an	tibody cream					
3 Betamethasone 17-valer		, ,				
Santoianni 2001	42	2.28 (3.03)	39	3.94 (3.03)	-	-0.54 [-0.99, -0.10]
4 Caffeine (topical) 10%, T						
Vali 2005	39	4.41 (4.83)	39	6.29 (4.71)		-0.39 [-0.84, 0.06]
5 Calcipotriene 0.005% oii		5% or 0.1% or 0.7%	5 or 1.4%, twic	e daily		
6 Dead Sea salts emollient Cheesbrough 1992	lotion, 30% 8	24 (11.1)	11	18.5 (7.5)		0.57 [-0.36, 1.51
0	0	21(11.1)		10.5 (7.5)		0.57 [-0.50, 1.51
7 Fish oil plus occlusion						
8 Herbal skin care (Dr Mic Maier 2004	haels cleansing gel, ointr 14	nent and skin condit -0.89 (0.15)	tioner), twice (10	daily -0.22 (0.29)		-2.96 [-4.19, -1.74
		-0.07 (0.13)	10	-0.22 (0.27)		-2.70 [- 1.17, -1.71]
9 Hexafluoro-1,25-dihydro	, ,					
10 Indigo naturalis 1.4% oi	ntment					
I I Kukui nut oil, twice dail	·					
Brown 2005	13	3.9 (3)	11	4 (2.7)		-0.03 [-0.84, 0.77]
12 Mahonia aquifoliu	, , ,					
Bernstein 2006	100	-3.39 (3.59)	100	-0.09 (4.85)	+	-0.77 [-1.06, -0.48]
13 Methotrexate gel						
Syed 2001b	30	2.2 (3.76)	30	8.2 (3.76)	+	-1.58 [-2.16, -0.99]
14 Mycophenolic acid oint	ment					
15 NG-monomethyl-L-arg	inine (L-NMMA) cream					
16 Nicotinamide 1.4%, twi	ce daily					
17 Oleum horwathiensis (Psoricur)					
18 Omega-3-polyunsatura	,					
					-10 -5 0 5	10
				Favours	other treatment Favours	s placebo
						(Continued

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Other treatment		Placebo		Std. Mean Difference	(Continued) Std. Difference
	N	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% CI
19 Platelet aggregation activ	vating factor (PAF)(Ro 24-	0238)				
20 Polymyxin B cream 200,	000 U/g					
21 PTH (1-34) in Novason	ne A liposomal cream, tv	vice daily				
22 Sirolimus (topical), 2.2%	for 6 wks, then 8% for a f	urther 6 wks				
23 Tacrolimus ointment						
24 Tar						
25 Tazarotene						
26 Theophylline 1% ointme	ent, twice daily					
Papakostantinou 2005	11	-0.5 (0.19)	11	-0.09 (0.04)		-2.87 [-4.13, -1.62]
					-10 -5 0 5 1	0

Favours other treatment Favours placebo

Analysis 6.4. Comparison 6 Other treatment versus placebo, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: 4 PAGI

Study or subgroup	Other treatment		Placebo		Std. Mean Difference	Std Mean Difference
,	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% C
2 Anti-IL-8 monoclona	% hydrophilic cream, thre I antibody cream alerate 21-acetate plus tre 42	. ,	cid 39	-1.69 (1.127)	-	-0.80 [-1.26, -0.35
6 Dead Sea salts emoll 7 Fish oil plus occlusior	6 ointment + nicotinamide ient lotion, 30%			,		
9 Hexafluoro-1,25-dihy	droxyvitamin D3, twice d	aily				
10 Indigo naturalis 1.4%	6 ointment					
I I Kukui nut oil, TD Brown 2005	13	1.7 (0.9)	П	1.7 (0.9)	-	0.0 [-0.80, 0.80
12 Mahonia aquifo	<i>lium</i> (Reli va™), twice o	daily				
13 Methotrexate gel						
14 Mycophenolic acid	ointment					
15 NG-monomethyl-L-	-arginine (L-NMMA) crea	n				
16 Nicotinamide 1.4%,	twice daily					
17 Oleum horwathiens	sis (Psoricur)					
18 Omega-3-polyunsat	urated fatty acids ointmer	nt				
19 Platelet aggregation	activating factor (PAF)(Rc	24-0238)				
20 Polymyxin B cream	200,000 U/g					
21 PTH (1-34) in Nova	asome A liposomal crea	m, twice daily				
22 Sirolimus (topical), 2	2.2% for 6 wks, then 8% fo	or a further 6 wks				
23 Tacrolimus ointmen	t					
24 Tar						
25 Tazarotene						
26 Theophylline 1% oir	ntment, twice daily					
					 10 -5 0 5	
					10 -5 0 5 her treatment Favours pla	10 icebo

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Analysis 6.5. Comparison 6 Other treatment versus placebo, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Other treatment		Placebo		Std. Mean Difference	Weight	Std. Mean Difference
study of subgroup	N	Mean(SD)	N	Mean(SD)	IV,Random,95% Cl	, reight	IV,Random,95% CI
I Aloe vera extract 0.5%	hydrophilic cream, th	ree times per day	/				
Syed 1996	30	2.2 (3.76)	30	8.2 (3.76)	-	100.0 %	-1.58 [-2.16, -0.99]
Subtotal (95% CI)	30		30		•	100.0 %	-1.58 [-2.16, -0.99]
Heterogeneity: not applica	able						
Test for overall effect: Z =	5.29 (P < 0.00001)						
2 Anti-IL-8 monoclonal ar	ntibody cream						
Jin 2001	45	-1.27 (1.01)	44	-0.73 (0.79)	+	100.0 %	-0.59 [-1.01, -0.16
Subtotal (95% CI)	45		44		•	100.0 %	-0.59 [-1.01, -0.16]
Heterogeneity: not applica	able						
Test for overall effect: Z =	= 2.72 (P = 0.0065)						
3 Betamethasone 17-vale	rate 21-acetate plus t	retinoin plus salic	ylic acid				
Santoianni 2001	42	-2.36 (0.69)	39	-1.79 (0.8)	-	100.0 %	-0.76 [-1.21, -0.31]
Subtotal (95% CI)	42		39		•	100.0 %	-0.76 [-1.21, -0.31]
Heterogeneity: not applica	able						
Test for overall effect: Z =	= 3.29 (P = 0.0010)						
4 Caffeine (topical) 10%,	TD						
Vali 2005	39	4.41 (4.83)	39	6.29 (4.71)	+	100.0 %	-0.39 [-0.84, 0.06]
Subtotal (95% CI)	39		39		•	100.0 %	-0.39 [-0.84, 0.06]
Heterogeneity: not applica	able						
Test for overall effect: Z =	= 1.71 (P = 0.088)						
5 Calcipotriene 0.005% oi	intment + nicotinami	de 0.05% or 0.1%	or 0.7% o	or 1.4%, twice daily	,		
Levine 2010 (P)	144	3.15 (1.94)	48	4.08 (1.93)	-	100.0 %	-0.48 [-0.81, -0.15]
Subtotal (95% CI)	144		48		•	100.0 %	-0.48 [-0.81, -0.15]
Heterogeneity: not applica	able						
Test for overall effect: Z =	= 2.84 (P = 0.0045)						
6 Dead Sea salts emollien	t lotion, 30%						
Cheesbrough 1992	8	24 (.)	11	18.5 (7.5)	•••	100.0 %	0.57 [-0.36, 1.51
Subtotal (95% CI)	8		11		•	100.0 %	0.57 [-0.36, 1.51]
Heterogeneity: not applica	able						
						1	
				-10		10	
				Favours other	treatment Favours pla	cebo	,

(Continued . . .)

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Other treatment	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	(Continued Std. Mean Difference IV,Random,95% CI
Test for overall effect: $Z =$	1.20 (P = 0.23)						
7 Fish oil plus occlusion Escobar 1992	25	1.74 (1.94)	25	3.81 (1.94)	-	100.0 %	-1.05 [-1.64, -0.46]
Subtotal (95% CI)	25		25		•	100.0 %	-1.05 [-1.64, -0.46]
Heterogeneity: not applical Test for overall effect: Z =							
8 Herbal skin care (Dr Mic Maier 2004	haels cleansing gel, 14	ointment and s -0.89 (0.15)	kin conditic 10	oner), twice daily -0.22 (0.29)		100.0 %	-2.96 [-4.19, -1.74]
Subtotal (95% CI) Heterogeneity: not applical Test for overall effect: Z =			10		•	100.0 %	-2.96 [-4.19, -1.74]
9 Hexafluoro-1,25-dihydro Durakovic 2001	xyvitamin D3 15	-3.3 (0.71)	15	-2.8 (0.86)	-	100.0 %	-0.62 [-1.35, 0.12]
Subtotal (95% CI)	15	-3.3 (0.71)	15	-2.0 (0.00)	•	100.0 %	-0.62 [-1.35, 0.12]
Heterogeneity: not applical Test for overall effect: Z =	ole		19			100.0 /0	[1:03,0:12]
10 Indigo naturalis 1.4% oir	ntment						
Lin 2007	10	3.7 (2)	10	7.2 (1.4)		22.9 %	-1.94 [-3.05, -0.84]
Lin 2008	34	-4.18 (1.22)	34	-1.68 (1.09)		77.1 %	-2.14 [-2.74, -1.53]
Subtotal (95% CI) Heterogeneity: Tau ² = 0.0; Test for overall effect: Z =		(P = 0.76); I ² =	44 0.0%		•	100.0 %	-2.09 [-2.62, -1.56]
I I Kukui nut oil, TD Brown 2005	13	1.7 (0.9)	11	1.7 (0.8)	-	100.0 %	0.0 [-0.80, 0.80]
Subtotal (95% CI)	13	(0)	11	(0.0)	•	100.0 %	0.0 [-0.80, 0.80]
Heterogeneity: not applical Test for overall effect: Z =	ble					10000 /0	
12 <i>Mahonia aquifoliun</i> Bernstein 2006	m (Reli va™), twice 100	daily -3.39 (3.59)	100	-0.09 (4.85)	+	100.0 %	-0.77 [-1.06, -0.48]
Subtotal (95% CI)	100	-5.57 (5.57)	100	-0.07 (1.03)	•		-0.77 [-1.06, -0.48]
Heterogeneity: not applical Test for overall effect: Z =	ole		100			100.0 /0	-0.77 [-1.00, -0.10]
13 Methotrexate gel Sutton 2001	41	-2.34 (1.2)	41	-1.78 (0.72)	-	51.8 %	-0.56 [-1.00, -0.12]
Syed 2001b	30	2.2 (3.76)	30	8.2 (3.76)		48.2 %	-1.58 [-2.16, -0.99]
Subtotal (95% CI) Heterogeneity: Tau ² = 0.44	71 I ; Chi ² = 7.37, df =	I (P = 0.0 I); I ²	71 =86%		•	100.0 %	-1.05 [-2.04, -0.06]
				-10 Favours other		10 acebo	(Continued

Study or subgroup C	Other treatment N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	(Continued Std. Mean Difference IV,Random,95% CI
Test for overall effect: $Z = 2.0$	07 (P = 0.038)						
14 Mycophenolic acid ointme Geilen 2000	ent 7	5.13 (0.73)	7	6.32 (0.81)		100.0 %	-1.44 [-2.67, -0.22]
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: $Z = 2$.			7		•	100.0 %	-1.44 [-2.67, -0.22]
15 NG-monomethyl-L-arginir	. ,						
Ormerod 2000	17	9.12 (3.55)	17	8.82 (4.22)	-	100.0 %	0.08 [-0.60, 0.75]
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: $Z = 0.2$			17		•	100.0 %	0.08 [-0.60, 0.75]
16 Nicotinamide 1.4%, twice Levine 2010 (P)	daily 48	3.69 (1.93)	48	4.08 (1.93)	-	100.0 %	-0.20 [-0.60, 0.20]
	48	5.07 (1.75)	48	1.00 (1.75)	Ţ		
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 0.9	e		40			100.0 %	-0.20 [-0.60, 0.20]
17 Oleum horwathiensis (Pso Lassus 1991	oricur) 19	-2.21 (1.69)	23	-2.17 (1.72)	=	100.0 %	-0.02 [-0.63, 0.58]
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: $Z = 0.0$			23		•	100.0 %	-0.02 [-0.63, 0.58]
18 Omega-3-polyunsaturated Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: not app	0	ent	0				Not estimable
 19 Platelet aggregation activa Wolska 1995 	ting factor (PAF)(F 40	,	40	-3.08 (0.73)	-	100.0 %	007 [050 027]
Subtotal (95% CI)	40	-3.13 (0.79)	40	-3.08 (0.73)	•	100.0 %	-0.07 [-0.50, 0.37] -0.07 [-0.50, 0.37]
Heterogeneity: not applicable Test for overall effect: $Z = 0.2$							
20 Polymyxin B cream 200,00 Stutz 1996	00 U/g 15	2.5 (1.5)	15	2.3 (1.5)	-	100.0 %	0.13 [-0.59, 0.85]
Subtotal (95% CI)	15		15		+	100.0 %	0.13 [-0.59, 0.85]
Heterogeneity: not applicable Test for overall effect: Z = 0.1							
21 PTH (1-34) in Novasome Holick 2003	A liposomal cre	am, twice daily -0.67 (0.23)	15	-0.18 (0.18)	-	100.0 %	-2.31 [-3.26, -1.36]
	13	, (0.20)	15	. ,		1	
				-10 Favours other		10 :ebo	(Continued

Study or subgroup	Other treatment		Placebo		Mean Difference	Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% Cl	-	IV,Random,95% CI
Subtotal (95% CI)	15		15		•	100.0 %	-2.31 [-3.26, -1.36]
Heterogeneity: not applicable Test for overall effect: $Z = 4$.							
22 Sirolimus (topical), 2.2% f	or 6 wks, then 8%	for a further 6 w	/ks				
Ormerod 2005	22	9.1 (4.8)	22	11.2 (5.8)		100.0 %	-0.39 [-0.98, 0.21]
Subtotal (95% CI) Heterogeneity: not applicabl Test for overall effect: Z = 1.			22		+	100.0 %	-0.39 [-0.98, 0.21]
23 Tacrolimus ointment Zonneveld 1998 (P)	24	4.7 (1.73)	23	4.6 (1.73)	-	100.0 %	0.06 [-0.52, 0.63]
Subtotal (95% CI) Heterogeneity: not applicabl Test for overall effect: Z = 0.			23		•	100.0 %	0.06 [-0.52, 0.63]
24 Tar							
Kanzler 1993	18	-3.25 (2.03)	18	-2.36 (1.86)	-	100.0 %	-0.45 [-1.11, 0.22]
Subtotal (95% CI) Heterogeneity: not applicabl Test for overall effect: Z = 1.			18		•	100.0 %	-0.45 [-1.11, 0.22]
25 Tazarotene							
Weinstein 1996	211	3.8 (1.73)	107	5.3 (1.73)	•	100.0 %	-0.86 [-1.11, -0.62]
Subtotal (95% CI) Heterogeneity: not applicabl Test for overall effect: Z = 7.			107		•	100.0 %	-0.86 [-1.11, -0.62]
26 Theophylline 1% ointmer	nt, twice daily						
Papakostantinou 2005	11	-0.5 (0.19)	11	-0.09 (0.04)		100.0 %	-2.87 [-4.13, -1.62]
Subtotal (95% CI) Heterogeneity: not applicabl Test for overall effect: Z = 4. Test for subgroup difference:	.48 (P < 0.00001)	F = 24 (P = 0.00	11), ² =81%		•	100.0 %	-2.87 [-4.13, -1.62]

Favours other treatment Favours placebo

Analysis 6.6. Comparison 6 Other treatment versus placebo, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: 6 Total withdrawals

Study or subgroup	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 CI
I Aloe vera extract 0.5% hydr	ophilic cream, three times	per day			
Syed 1996	0/30	0/30	+	100.0 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	30	30	+	100.0 %	0.0 [-0.06, 0.06]
Total events: 0 (Other treatme	ent), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
2 Anti-IL-8 monoclonal antibo	dy cream				
Jin 2001	3/48	4/48		100.0 %	-0.02 [-0.12, 0.08]
Subtotal (95% CI)	48	48	+	100.0 %	-0.02 [-0.12, 0.08]
Total events: 3 (Other treatme	ent), 4 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.39$	9 (P = 0.69)				
3 Betamethasone 17-valerate 2	21-acetate plus tretinoin p	lus salicylic acid			
Santoianni 2001	2/44	2/41	-	100.0 %	0.00 [-0.09, 0.09]
Subtotal (95% CI)	44	41	+	100.0 %	0.00 [-0.09, 0.09]
Total events: 2 (Other treatme	ent), 2 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.07$	7 (P = 0.94)				
4 Caffeine (topical) 10%, TD					
Vali 2005	3/39	3/39		100.0 %	0.0 [-0.12, 0.12]
Subtotal (95% CI)	39	39	+	100.0 %	0.0 [-0.12, 0.12]
Total events: 3 (Other treatme	ent), 3 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
5 Calcipotriene 0.005% ointme	ent + nicotinamide 0.05%	or 0.1% or 0.7% or 1	.4%, twice daily		
Levine 2010 (P)	5/144	0/48	+	100.0 %	0.03 [-0.01, 0.08]
Subtotal (95% CI)	144	48	•	100.0 %	0.03 [-0.01, 0.08]
Total events: 5 (Other treatme	nt), 0 (Placebo)				
Heterogeneity: not applicable					
	2 (P = 0.10)				
Test for overall effect: $Z = 1.62$					
lest for overall effect: $\angle = 1.6$. 6 Dead Sea salts emollient loti	on				

			Risk		Risk
Study or subgroup	Other treatment	Placebo	Difference M-	Weight	Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
Subtotal (95% CI)	12	12	-	100.0 %	0.25 [-0.06, 0.56]
otal events: 4 (Other treatm	nent), I (Placebo)				
Heterogeneity: not applicable	2				
est for overall effect: $Z = 1.5$	58 (P = 0.11)				
7 Fish oil plus occlusion					
Escobar 1992	0/25	0/25		100.0 %	0.0 [-0.07, 0.07]
Subtotal (95% CI)	25	25	+	100.0 %	0.0 [-0.07, 0.07]
Total events: 0 (Other treatm	nent), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
3 Herbal skin care (Dr Micha	els cleansing gel, ointmer	t and skin conditioner), twice daily		
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatm	nent), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not ap					
Hexafluoro-1,25-dihydroxy		0/15		100.0.0/	
Durakovic 2001	0/15	0/15		100.0 %	0.0 [-0.12, 0.12]
Subtotal (95% CI)	15	15	+	100.0 %	0.0 [-0.12, 0.12]
Total events: 0 (Other treatm	, , ,				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
10 Indigo naturalis 1.4% ointr	ment				
Lin 2007	4/14	4/14		20.1 %	0.0 [-0.33, 0.33]
Lin 2008	8/42	8/42	-	79.9 %	0.0 [-0.17, 0.17]
Subtotal (95% CI)	56	56	+	100.0 %	0.0 [-0.15, 0.15]
Fotal events: 12 (Other treat	ment), 12 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$; C); I ² =0.0%			
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
I I Kukui nut oil, TD					
Brown 2005	2/15	4/15	— <u>—</u>	100.0 %	-0.13 [-0.42, 0.15]
Subtotal (95% CI)	15	15	-	100.0 %	-0.13 [-0.42, 0.15]
Total events: 2 (Other treatm					
-leterogeneity: not applicable	, , ,				
Test for overall effect: Z = 0.9	93 (P = 0.35)				
A Mahani a and filing					
2 <i>Mahonia aquifolium</i> Bernstein 2006	(Reli Va'''), twice daily 3/100	26/100		100.0 %	-0.23 [-0.32, -0.14]
Subtotal (95% CI)	100	100	•	100.0 %	-0.23 [-0.32, -0.14]
Total events: 3 (Other treatm					
Heterogeneity: not applicable	2				
			<u> </u>		
			-1 -0.5 0 0.5 1		

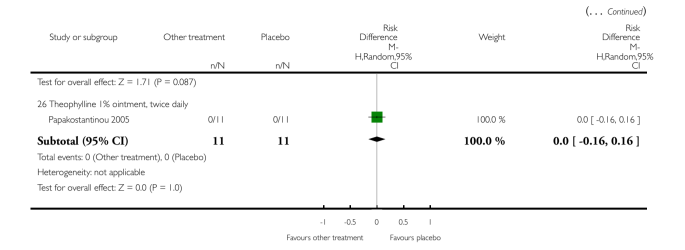
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Study or subgroup	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
est for overall effect: $Z = 4$.	89 (P < 0.00001)				
3 Methotrexate gel					
Syed 2001b	0/30	0/30	-	100.0 %	0.0 [-0.06, 0.06]
Subtotal (95% CI) Total events: 0 (Other treatme Heterogeneity: not applicable Test for overall effect: Z = 0.	e	30	•	100.0 %	0.0 [-0.06, 0.06]
4 Mycophenolic acid ointm				100.0.0/	
Geilen 2000	0/7	0/7	–	100.0 %	0.0 [-0.24, 0.24]
Subtotal (95% CI) Fotal events: 0 (Other treatn Heterogeneity: not applicable Fest for overall effect: Z = 0.	e	7		100.0 %	0.0 [-0.24, 0.24]
15 NG-monomethyl-L-argini Ormerod 2000	ine (L-NMMA) cream 0/17	0/17	-	100.0 %	0.0 [-0.11, 0.11]
Subtotal (95% CI) Total events: 0 (Other treatn Heterogeneity: not applicable Test for overall effect: Z = 0.	e	17	+	100.0 %	0.0 [-0.11, 0.11]
6 Nicotinamide 1.4%, twice Levine 2010 (P)	e daily 1/48	0/48	-	100.0 %	0.02 [-0.04, 0.08]
Subtotal (95% CI) otal events: I (Other treatmeterogeneity: not applicable est for overall effect: Z = 0.	e	48	+	100.0 %	0.02 [-0.04, 0.08]
7 Oleum horwathiensis Lassus 1991	6/25	2/25		100.0 %	
Subtotal (95% CI) Fotal events: 6 (Other treatme Heterogeneity: not applicable Fest for overall effect: Z = 1.	25 nent), 2 (Placebo) e	25	-	100.0 %	0.16 [-0.04, 0.36]
8 Omega-3-polyunsaturate Henneicke-v. Z. 1993	d fatty acids ointment 21/73	21/73	-	100.0 %	0.0 [-0.15, 0.15]
Subtotal (95% CI)	73	73	Ţ	100.0 %	
bubtotal (95% CI) otal events: 21 (Other treat leterogeneity: not applicable est for overall effect: $Z = 0$.	tment), 21 (Placebo) e	/3		100.0 %	0.0 [-0.15, 0.15]
9 Platelet aggregation activa	ating factor (PAF)(Ro 24-02.	38)			

(Continued . . .)

(... Continued)

Risk Difference M-	Weight	Risk Difference M-	Placebo	Other treatment	Study or subgroup
H,Random,S C		H,Random,95% Cl	n/N	n/N	
0.0 [-0.16, 0.16]	100.0 %	-	12/52	12/52	Wolska 1995
0.0 [-0.16, 0.16]	100.0 %	+	52	, , , ,	Subtotal (95% CI) Total events: 12 (Other treatr Heterogeneity: not applicable Test for overall effect: Z = 0.0
0.0 [-0.24, 0.24]	100.0 %		2/15	10 U/g 2/15	20 Polymyxin B cream 200,00 Stutz 1996
0.0 [-0.24, 0.24]	100.0 %	⊥ I	15	15	Subtotal (95% CI)
0.0 [-0.24, 0.24]	100.0 /0		19	ent), 2 (Placebo)	Total events: 2 (Other treatm Heterogeneity: not applicable Test for overall effect: Z = 0.0
0.0 [-0.12, 0.12	100.0 %	-	e daily 0/15	A liposomal cream, twice 0/15	21 PTH (1-34) in Novasome Holick 2003
0.0 [-0.12, 0.12]	100.0 %	•	15	15	Subtotal (95% CI)
Not estimable			0	0 (P = 1.0) 0 ent), 0 (Placebo)	Total events: 0 (Other treatm Heterogeneity: not applicable Test for overall effect: Z = 0.0 22 Sirolimus (topical) Subtotal (95% CI) Total events: 0 (Other treatm
Not estimable			0	olicable 0 ent), 0 (Placebo)	Heterogeneity: not applicable Fest for overall effect: not app 23 Tacrolimus ointment Subtotal (95% CI) Fotal events: 0 (Other treatm Heterogeneity: not applicable
Not estimable			0	olicable 0	Test for overall effect: not app 24 Tar Subtotal (95% CI)
			,	ent), 0 (Placebo)	Fotal events: 0 (Other treatm Heterogeneity: not applicable Test for overall effect: not app
0.00 [-0.10, 0.10]	21.5 %	+	27/108	55/216	25 Tazarotene Weinstein 1996
0.05 [0.00, 0.10	78.5 %	-	125/443	286/860	Weinstein 2003
0.04 [-0.01, 0.09]	100.0 %	•	551	1076 tment), I52 (Placebo)	Subtotal (95% CI) Fotal events: 341 (Other treat Heterogeneity: Tau ² = 0.0; Ch



Analysis 6.7. Comparison 6 Other treatment versus placebo, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,9 Cl	5%	H,Random,95% Cl
I Aloe vera extract 0.5% hy	drophilic cream, three times	per day			
Syed 1996	0/30	0/30	-	100.0 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	30	30	+	100.0 %	0.0 [-0.06, 0.06]
Total events: 0 (Other treatr	ment), 0 (Placebo)				
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 0$.	.0 (P = 1.0)				
2 Anti-IL-8 monoclonal antib	oody cream				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatr	ment), 0 (Placebo)				
Heterogeneity: not applicable	e				
Test for overall effect: not ap	plicable				
				1 1	
			-1 -0.5 0 0).5 I	
		Favou	rs other treatment Fav	ours placebo	
					(Continued)

(... Continued)

			D'-L		(Continued
Study or subgroup	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
3 Betamethasone 17-valerate 21	-acetate plus tretinoin p	us salicylic acid			
Santoianni 2001	1/44	1/41	-	100.0 %	0.00 [-0.07, 0.06]
Subtotal (95% CI)	44	41	+	100.0 %	0.00 [-0.07, 0.06]
Total events: (Other treatment	:), I (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.05$ (P = 0.96)				
4 Caffeine (topical) 10%, TD					
Vali 2005	0/39	0/39	+	100.0 %	0.0 [-0.05, 0.05]
Subtotal (95% CI)	39	39	•	100.0 %	0.0 [-0.05, 0.05]
Total events: 0 (Other treatment	:), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ (P	= 1.0)				
5 Calcipotriene 0.005% ointmen	t + nicotinamide 0.05%	or 0.1% or 0.7% or 1.4	1%, twice daily		
Levine 2010 (P)	0/144	0/48	-	100.0 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	144	48	•	100.0 %	0.0 [-0.03, 0.03]
Total events: 0 (Other treatment		10		10010 /0	
Heterogeneity: not applicable	,, - (
Test for overall effect: Z = 0.0 (P	= 1.0)				
6 Dead Sea salts emollient lotion					
Cheesbrough 1992	2/12	1/12		100.0 %	0.08 [-0.18, 0.35]
Subtotal (95% CI)	12	12	-	100.0 %	0.08 [-0.18, 0.35]
Total events: 2 (Other treatment		12		100.0 /0	0.00 [0.10, 0.55]
Heterogeneity: not applicable	,, (
Test for overall effect: Z = 0.62 (P = 0.53)				
7 Fish oil plus occlusion					
Escobar 1992	0/25	0/25	-	100.0 %	0.0 [-0.07, 0.07]
Subtotal (95% CI)	25	25	•	100.0 %	0.0 [-0.07, 0.07]
Total events: 0 (Other treatment	-	29		100.0 /0	
Heterogeneity: not applicable), o (1 lacebo)				
Test for overall effect: $Z = 0.0$ (P	= .0)				
B Herbal skin care (Dr Michaels	,	and skin conditioner)	twice daily		
Subtotal (95% CI)	0	0	, twice daily		Not estimable
Total events: 0 (Other treatment		Ū			
Heterogeneity: not applicable), o (i iaccoo)				
Test for overall effect: not applica	able				
יי אראי Hexafluoro-1,25-dihydroxyvita					
Durakovic 2001	0/15	0/15	H	100.0 %	0.0 [-0.12, 0.12]
Subtotal (95% CI)	15	15	+	100.0 %	0.0 [-0.12, 0.12]
Total events: 0 (Other treatment		-2			
Heterogeneity: not applicable	,				
			-1 -0.5 0 0.5 1		
		Favours ot	her treatment Favours placeb	0	
					(Continued

Risl Difference	Weight	Risk Difference	Placebo	Other treatment	Study or subgroup
M H,Random, C	0	M- H,Random,95% Cl	n/N	n/N	, , ,
					Test for overall effect: $Z = 0.0$
				nent	10 Indigo naturalis 1.4% ointn
0.0 [-0.13, 0.13	11.1 %		0/14	0/14	Lin 2007
0.0 [-0.05, 0.05	88.9 %	-	0/42	0/42	Lin 2008
0.0 [-0.04, 0.04]	100.0 %	•	56	56	Subtotal (95% CI)
			$ ^2 = 0.0\%$	$hi^2 = 0.0, df = 1 (P = 1.00);$	Total events: 0 (Other treatm Heterogeneity: Tau ² = 0.0; Cł Test for overall effect: Z = 0.0
					l I Kukui nut oil, TD
0.0 [-0.12, 0.12	100.0 %	=	0/15	0/15	Brown 2005
0.0 [-0.12, 0.12]	100.0 %	•	15	, , ,	Subtotal (95% CI) Total events: 0 (Other treatm Heterogeneity: not applicable Test for overall effect: Z = 0.0
Not estimable			0	0 ent), 0 (Placebo)	12 <i>Mahonia aquifolium</i> Subtotal (95% CI) Total events: 0 (Other treatm Heterogeneity: not applicable Test for overall effect: not app
0.0 [-0.06, 0.06	100.0 %		0/30	0/30	13 Methotrexate gel Syed 2001b
0.0 [-0.06, 0.06]	100.0 %	•	30	, , ,	Subtotal (95% CI) Total events: 0 (Other treatme Heterogeneity: not applicable Test for overall effect: Z = 0.0
				nt	14 Mycophenolic acid ointme
0.0 [-0.24, 0.24	100.0 %		0/7	0/7	Geilen 2000
0.0 [-0.24, 0.24]	100.0 %	-	7	, , ,	Subtotal (95% CI) Fotal events: 0 (Other treatm Heterogeneity: not applicable Fest for overall effect: Z = 0.0
0.0 [-0.11, 0.11]	100.0 %	-	0/17	ne (L-NMMA) cream 0/17	5 NG-monomethyl-L-arginir Ormerod 2000
0.0 [-0.11, 0.11]	100.0 %	+	17	, , ,	Subtotal (95% CI) Total events: 0 (Other treatme Heterogeneity: not applicable Test for overall effect: Z = 0.0

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Risk Difference M-	Weight	Risk Difference M-	Placebo	Other treatment	Study or subgroup
H,Random, C		H,Random,95% Cl	n/N	n/N	
				daily	6 Nicotinamide 1.4%, twice o
0.0 [-0.04, 0.04]	100.0 %	-	0/48	0/48	Levine 2010 (P)
0.0 [-0.04, 0.04]	100.0 %	•	48	48	Subtotal (95% CI)
				, , ,	Total events: 0 (Other treatme Heterogeneity: not applicable Test for overall effect: Z = 0.0
0.0 [-0.07, 0.07]	100.0 %	-	0/25	0/25	17 Oleum horwathiensis Lassus 1991
0.0 [-0.07, 0.07]	100.0 %	•	25	25 ent), 0 (Placebo)	Subtotal (95% CI) Total events: 0 (Other treatme
					Heterogeneity: not applicable Test for overall effect: $Z = 0.0$
0.0 [-0.03, 0.03]	100.0 %	-	0/73	fatty acids ointment 0/73	18 Omega-3-polyunsaturated Henneicke-v. Z. 1993
0.0 [-0.03, 0.03]	100.0 %	•	73	, , ,	Subtotal (95% CI) Total events: 0 (Other treatme Heterogeneity: not applicable Test for overall effect: Z = 0.0
Not estimable			³⁾ 0	0 ent), 0 (Placebo)	19 Platelet aggregation activati Subtotal (95% CI) Total events: 0 (Other treatme Heterogeneity: not applicable Test for overall effect: not app
Not estimable			0	0 ent), 0 (Placebo)	20 Polymyxin B cream 200,000 Subtotal (95% CI) Total events: 0 (Other treatme Heterogeneity: not applicable Fest for overall effect: not app
			daily	A liposomal cream, twice	21 PTH (1-34) in Novasome /
0.0 [-0.12, 0.12]	100.0 %	=	0/15	0/15	Holick 2003
0.0 [-0.12, 0.12]	100.0 %	+	15	, , ,	Subtotal (95% CI) Total events: 0 (Other treatme Heterogeneity: not applicable Test for overall effect: Z = 0.0
Not estimable			0		22 Sirolimus (topical) Subtotal (95% CI) Total events: 0 (Other treatme Heterogeneity: not applicable Test for overall effect: not app

(Continued . . .)

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Study or subgroup	Other treatment	Placebo	Risk Difference	Weight	Risk Difference
			M- H,Random,95%		M- H,Random,959
	n/N	n/N	Ċ		ĊI
23 Tacrolimus ointment					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatmer	nt), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applie	cable				
24 Tar					
Subtotal (95% CI)	0	0			Not estimable
Fotal events: 0 (Other treatmer	nt), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applie	cable				
25 Tazarotene					
Weinstein 1996	24/216	3/108	-	23.2 %	0.08 [0.03, 0.14]
Weinstein 2003	97/860	20/443	-	76.8 %	0.07 [0.04, 0.10]
Subtotal (95% CI)	1076	551	•	100.0 %	0.07 [0.05, 0.10]
Fotal events: 121 (Other treatm	nent), 23 (Placebo)				
Heterogeneity: Tau ² = 0.0; Chi ²	² = 0.27, df = 1 (P = 0.60); I ² =0.0%			
Fest for overall effect: Z = 5.56	(P < 0.00001)				
26 Theophylline 1% ointment, t	wice daily				
Papakostantinou 2005	0/11	0/11		100.0 %	0.0 [-0.16, 0.16]
Subtotal (95% CI)	11	11	+	100.0 %	0.0 [-0.16, 0.16]
Total events: 0 (Other treatmer	nt), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 0.0 (P = 1.0				

Favours other treatment Favours placebo

Analysis 6.8. Comparison 6 Other treatment versus placebo, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Aloe vera extract 0.5% hyd	frophilic cream, three times	per day			
Syed 1996	0/30	0/30	-	100.0 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	30	30	+	100.0 %	0.0 [-0.06, 0.06]
Total events: 0 (Other treatm	nent), 0 (Placebo)				
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
2 Anti-IL-8 monoclonal antib	ody cream				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatm	, , , ,				
Heterogeneity: not applicable					
Test for overall effect: not app					
3 Betamethasone 17-valerate Santoianni 2001	e 21-acetate plus tretinoin p 1/44	lus salicylic acid 0/41	_	100.0 %	0.02 [-0.04, 0.08]
Santolanni 2001	1/44	0/41	—	100.0 %	0.02 [-0.04, 0.08]
Subtotal (95% CI)	44	41	•	100.0 %	0.02 [-0.04, 0.08]
Total events: I (Other treatm	nent), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.7$	()				
4 Caffeine (topical) 10%, TD		0/20	-		
Vali 2005	0/39	0/39		100.0 %	0.0 [-0.05, 0.05]
Subtotal (95% CI)	39	39	•	100.0 %	0.0 [-0.05, 0.05]
Total events: 0 (Other treatm	, , ,				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	· · · ·	0.10/ 0.70/ 1	10(
5 Calcipotriene 0.005% ointn			.4%, twice daily	100.0 %	
Levine 2010 (P)	0/144	0/48	—	100.0 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	144	48	•	100.0 %	0.0 [-0.03, 0.03]
Total events: 0 (Other treatm	, , , ,				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	· · · ·				
6 Dead Sea salts emollient lo		0/12			
Cheesbrough 1992	1/12	0/12		100.0 %	0.08 [-0.12, 0.29]
Subtotal (95% CI)	12	12	-	100.0 %	0.08 [-0.12, 0.29]
			-1 -0.5 0 0.5 1		
		Favours of	ther treatment Favours placeb	00	/ <u>-</u>
					(Continued

Study or subgroup	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random, C
Total events: (Other treatmer	nt), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.81$	(P = 0.42)				
7 Fish oil plus occlusion					
Escobar 1992	0/25	0/25	—	100.0 %	0.0 [-0.07, 0.07]
Subtotal (95% CI)	25	25	+	100.0 %	0.0 [-0.07, 0.07]
Total events: 0 (Other treatmer	nt), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ (P = 1.0)				
8 Herbal skin care (Dr Michaels	cleansing gel, ointment	t and skin conditioner), twice daily		
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatmer	nt), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applie					
9 Hexafluoro-1,25-dihydroxyvit		0/15		100.0.0/	
Durakovic 2001	0/15	0/15	-	100.0 %	0.0 [-0.12, 0.12]
Subtotal (95% CI)	15	15	+	100.0 %	0.0 [-0.12, 0.12]
Total events: 0 (Other treatmer	nt), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ (P = 1.0)				
10 Indigo naturalis 1.4% ointme	nt				
Lin 2007	0/14	0/14		100.0 %	0.0 [-0. 3, 0. 3
Subtotal (95% CI)	14	14	+	100.0 %	0.0 [-0.13, 0.13]
Total events: 0 (Other treatmer	nt), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ (P = 1.0)				
I I Kukui nut oil, TD					
Brown 2005	0/15	0/15		100.0 %	0.0 [-0.12, 0.12
Subtotal (95% CI)	15	15	•	100.0 %	0.0 [-0.12, 0.12]
Total events: 0 (Other treatmer	-	19		100.0 /0	0.0 [-0.12, 0.12]
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$ (P = 1.0)				
12 <i>Mahonia aquifolium</i> (R Subtotal (95% CI)	leli va™), twice daily 0	0			Not estimable
Total events: 0 (Other treatmer		U			inot estimable
Heterogeneity: not applicable	1, U (1 IACEUU)				
Test for overall effect: not applic	cable				
13 Methotrexate gel	0/20	0/20		100.0.9/	
Syed 2001b	0/30	0/30		100.0 %	0.0 [-0.06, 0.06]

(... Continued)

Study or subgroup	Other treatment	Placebo	Risk Difference	Weight Diffe		
,8,			M- H,Random,95%		H,Random,9	
Subtotal (95% CI)	n/N 30	n/N 30	CI	100.0 %	0.0 [-0.06, 0.06]	
Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: Z = 0.0 (F	t), 0 (Placebo)	90		100.0 /5	0.0 [-0.00, 0.00]	
14 Mycophenolic acid ointment Geilen 2000	0/7	0/7		100.0 %	0.0 [-0.24, 0.24]	
Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: Z = 0.0 (F		7	-	100.0 %	0.0 [-0.24, 0.24]	
15 NG-monomethyl-L-arginine Ormerod 2000	(L-NMMA) cream 0/17	0/17	=	100.0 %	0.0 [-0.11, 0.11]	
Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f		17	•	100.0 %	0.0 [-0.11, 0.11]	
16 Nicotinamide 1.4%, twice dai Levine 2010 (P)	ily 0/48	0/48	-	100.0 %	0.0 [-0.04, 0.04]	
Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f		48	ł	100.0 %	0.0 [-0.04, 0.04]	
17 Oleum horwathiensis Lassus 1991	0/25	0/25		100.0 %		
				100.0 %	0.0 [-0.07, 0.07]	
Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f	P = 1.0)	25		100.0 %	0.0 [-0.07, 0.07]	
18 Omega-3-polyunsaturated fa Henneicke-v. Z. 1993	0/73	0/73	•	100.0 %	0.0 [-0.03, 0.03]	
Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f		73	ł	100.0 %	0.0 [-0.03, 0.03]	
19 Platelet aggregation activating Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: not applic	0 t), 0 (Placebo)	³⁸⁾ 0			Not estimable	

(Continued . . .)

Study or subgroup	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
20 Polymyxin B cream 200,000 Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: not applic	0 t), 0 (Placebo)	0			Not estimable
21 PTH (1-34) in Novasome A Holick 2003	liposomal cream, twice 0/15	e daily 0/15	_	100.0 %	0.0 [-0.12, 0.12]
Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: Z = 0.0 (f	15 t), 0 (Placebo)	15	•	100.0 %	0.0 [-0.12, 0.12]
22 Sirolimus (topical) Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: not applic		0			Not estimable
23 Tacrolimus ointment Subtotal (95% CI) Total events: 0 (Other treatmen Heterogeneity: not applicable Test for overall effect: not applic	0 t), 0 (Placebo)	0			Not estimable
24 Tar Subtotal (95% CI) Fotal events: 0 (Other treatmen Heterogeneity: not applicable Fest for overall effect: not applic		0			Not estimable
25 Tazarotene Weinstein 1996	9/216	6/108	Ļ	21.6 %	-0.01 [-0.06, 0.04]
Weinstein 2003	40/860	28/443	•	78.4 %	-0.02 [-0.04, 0.01]
Subtotal (95% CI) Total events: 49 (Other treatme Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 1.34	= 0.01, df = 1 (P = 0.92)	551 2); I ² =0.0%	•	100.0 %	-0.02 [-0.04, 0.01]
26 Theophylline 1% ointment, tv Papakostantinou 2005	wice daily 0/11	0/11	-	100.0 %	0.0 [-0.16, 0.16]
Subtotal (95% CI) Fotal events: 0 (Other treatmen Heterogeneity: not applicable Fest for overall effect: Z = 0.0 (F	11 t), 0 (Placebo)	11	-	100.0 %	0.0 [-0.16, 0.16]

Analysis 6.9. Comparison 6 Other treatment versus placebo, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: 9 Adverse events (local)

 I Aloe vera extract 0.5% hydrophil Syed 1996 Subtotal (95% CI) Total events: 0 (Other treatment), 6 Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 2 Anti-IL-8 monoclonal antibody cr Jin 2001 Subtotal (95% CI) Total events: 5 (Other treatment), 4 Heterogeneity: not applicable 	0/30 30 D (Placebo) I.0) eam 5/46 46	0/30 30 4/46	M- H,Random,95% Cl	100.0 % 100.0 %	M- H,Random,9 CI 0.0 [-0.06, 0.06] 0.0 [-0.06, 0.06]
Syed 1996 Subtotal (95% CI) Total events: 0 (Other treatment), 0 Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 2 Anti-IL-8 monoclonal antibody cr Jin 2001 Subtotal (95% CI) Total events: 5 (Other treatment), 4	0/30 30 D (Placebo) I.0) eam 5/46 46	0/30 30 4/46	•	100.0 %	
Subtotal (95% CI) Total events: 0 (Other treatment), (Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 2 Anti-IL-8 monoclonal antibody cr Jin 2001 Subtotal (95% CI) Total events: 5 (Other treatment),	30 D (Placebo) I.0) eam 5/46 46	30 4/46	•	100.0 %	
Total events: 0 (Other treatment), 6 Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 2 Anti-IL-8 monoclonal antibody cr Jin 2001 Subtotal (95% CI) Total events: 5 (Other treatment), 4	D (Placebo) I.0) eam 5/46 46	4/46	•		0.0 [-0.06, 0.06]
Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 2 Anti-IL-8 monoclonal antibody cr Jin 2001 Subtotal (95% CI) Total events: 5 (Other treatment), 4	1.0) eam 5/46 46		-		
Test for overall effect: Z = 0.0 (P = 2 Anti-IL-8 monoclonal antibody cr Jin 2001 Subtotal (95% CI) Total events: 5 (Other treatment), 4	eam 5/46 46		-		
2 Anti-IL-8 monoclonal antibody cr Jin 2001 Subtotal (95% CI) Total events: 5 (Other treatment), 4	eam 5/46 46		-		
Jin 2001 Subtotal (95% CI) Total events: 5 (Other treatment), 4	5/46 46		-		
Subtotal (95% CI) Total events: 5 (Other treatment), 4	46				
Total events: 5 (Other treatment), 4				100.0 %	0.02 [-0.10, 0.14]
· · · · · · · · · · · · · · · · · · ·		46	+	100.0 %	0.02 [-0.10, 0.14]
Heterogeneity: not applicable	4 (Placebo)				
Test for overall effect: $Z = 0.35$ (P =	= 0.73)				
3 Betamethasone 17-valerate 21-ad	etate plus tretinoin pl	lus salicylic acid			
Santoianni 2001	1/44	1/41		100.0 %	0.00 [-0.07, 0.06]
Subtotal (95% CI)	44	41	•	100.0 %	0.00 [-0.07, 0.06]
Total events: (Other treatment),	l (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 0.05 (P =	= 0.96)				
4 Caffeine (topical) 10%, TD					
Vali 2005	2/39	0/39		100.0 %	0.05 [-0.03, 0.13]
Subtotal (95% CI)	39	39	◆	100.0 %	0.05 [-0.03, 0.13]
Total events: 2 (Other treatment), () (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: Z = 1.22 (P =	= 0.22)				
5 Calcipotriene 0.005% ointment +	nicotinamide 0.05%	or 0.1% or 0.7% or 1.4	1%, twice daily		
Levine 2010 (P)	51/144	11/48		100.0 %	0.13 [-0.02, 0.27]

(Continued . . .)

			Risk		Risk
Study or subgroup	Other treatment	Placebo	Difference M-	Weight	Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
Subtotal (95% CI)	144	48	•	100.0 %	0.13 [-0.02, 0.27]
Total events: 51 (Other treat Heterogeneity: not applicabl Test for overall effect: $Z = 1$ 6 Dead Sea salts emollient lo	le .72 (P = 0.085)	1/12		100.0 %	
Cheesbrough 1992				100.0 %	0.08 [-0.18, 0.35]
Subtotal (95% CI) Total events: 2 (Other treatr Heterogeneity: not applicabl Test for overall effect: Z = 0 7 Fish oil plus occlusion Escobar 1992	le	0/25		100.0 %	0.08 [-0.18, 0.35]
		25	I		0.04 [-0.06, 0.14]
Subtotal (95% CI) Total events: 1 (Other treatr Heterogeneity: not applicabl Test for overall effect: Z = 0 8 Herbal skin care (Dr Mich:	le .75 (P = 0.45)	-		100.0 %	0.01 [-0.00, 0.14]
Maier 2004	aeis cleansing gei, ointmen 3/14	3/10		100.0 %	-0.09 [-0.44, 0.27]
Subtotal (95% CI) Total events: 3 (Other treatr Heterogeneity: not applicabl Test for overall effect: Z = 0 9 Hexafluoro-1,25-dihydrox	le .47 (P = 0.64)	10	-	100.0 %	-0.09 [-0.44, 0.27]
Durakovic 2001	2/15	0/15		100.0 %	0.13 [-0.06, 0.33]
Subtotal (95% CI) Total events: 2 (Other treatr Heterogeneity: not applicabl Test for overall effect: Z = 1 10 Indigo naturalis 1.4% oint	le .32 (P = 0.19)	15	-	100.0 %	0.13 [-0.06, 0.33]
Lin 2007	0/10	0/10	-	9.3 %	0.0 [-0.17, 0.17]
Lin 2008	0/34	0/34	•	90.7 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	44	44	+	100.0 %	0.0 [-0.05, 0.05]
Total events: 0 (Other treatr Heterogeneity: Tau ² = 0.0; C Test for overall effect: $Z = 0$	$Chi^2 = 0.0, df = 1 (P = 1.00)$); I ² =0.0%			
I I Kukui nut oil, TD Brown 2005	0/15	0/15	-	100.0 %	0.0 [-0.12, 0.12]
Subtotal (95% CI)	15	15	+	100.0 %	0.0 [-0.12, 0.12]
Total events: 0 (Other treatr	ment), 0 (Placebo)				
			-1 -0.5 0 0.5 1		
		Favours ot	her treatment Favours placeb	0	(Continued
					(Conunued

Study or subgroup C	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P =	1.0)				
12 <i>Mahonia aquifolium</i> (Reli Bernstein 2006	va™), twice daily I/100	3/100		100.0 %	-0.02 [-0.06, 0.02]
Subtotal (95% CI) Total events: 1 (Other treatment), 3 Heterogeneity: not applicable Test for overall effect: Z = 1.01 (P =	· · · ·	100	•	100.0 %	-0.02 [-0.06, 0.02]
13 Methotrexate gel Syed 2001b	0/30	0/30		100.0 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	30	30	•	100.0 %	0.0 [-0.06, 0.06]
Total events: 0 (Other treatment), 0 Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P =) (Placebo)	50		100.0 /0	0.0 [-0.00, 0.00]
14 Mycophenolic acid ointment Geilen 2000	0/7	0/7		100.0 %	0.0 [-0.24, 0.24]
Subtotal (95% CI) Total events: 0 (Other treatment), C Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P =	. ,	7	-	100.0 %	0.0 [-0.24, 0.24]
15 NG-monomethyl-L-arginine (L-N Ormerod 2000	NMMA) cream 0/17	0/17	-	100.0 %	0.0 [-0.11, 0.11]
Subtotal (95% CI) Total events: 0 (Other treatment), C Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P =	. ,	17	-	100.0 %	0.0 [-0.11, 0.11]
16 Nicotinamide 1.4%, twice daily Levine 2010 (P)	16/48	11/48	-	100.0 %	0.10 [-0.07, 0.28]
Subtotal (95% CI) Total events: 16 (Other treatment), Heterogeneity: not applicable Test for overall effect: Z = 1.14 (P =		48	-	100.0 %	0.10 [-0.07, 0.28]
17 Oleum horwathiensis Lassus 1991	1/25	0/25		100.0 %	0.04 [-0.06, 0.14]
Subtotal (95% CI) Fotal events: 1 (Other treatment), C Heterogeneity: not applicable Fest for overall effect: Z = 0.75 (P =	· · · ·	25	•	100.0 %	0.04 [-0.06, 0.14]

(Continued . . .)

					(Continued
Study or subgroup	Other treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
8 Omega-3-polyunsaturated fatt;	y acids ointment				
Henneicke-v. Z. 1993	1/73	0/73	-	100.0 %	0.01 [-0.02, 0.05]
Subtotal (95% CI) Fotal events: I (Other treatment), Heterogeneity: not applicable Fest for overall effect: Z = 0.72 (P	. ,	73	•	100.0 %	0.01 [-0.02, 0.05]
9 Platelet aggregation activating f Wolska 1995	actor (PAF)(Ro 24-023 23/52	8) 23/52	-	100.0 %	0.0 [-0.19, 0.19]
Subtotal (95% CI) Total events: 23 (Other treatment Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P =	= 1.0)	52	-	100.0 %	0.0 [-0.19, 0.19]
20 Polymyxin B cream 200,000 U/ Subtotal (95% CI) Fotal events: 0 (Other treatment), Heterogeneity: not applicable Fest for overall effect: not applicab	0 0 (Placebo)	0			Not estimable
21 PTH (1-34) in Novasome A Holick 2003	liposomal cream, twice 0/15	daily 0/15	=	100.0 %	0.0 [-0.12, 0.12]
Subtotal (95% CI) Total events: 0 (Other treatment), Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P :		15	+	100.0 %	0.0 [-0.12, 0.12]
22 Sirolimus (topical) Subtotal (95% CI) Fotal events: 0 (Other treatment), Heterogeneity: not applicable Fest for overall effect: not applicab		0			Not estimable
23 Tacrolimus ointment Subtotal (95% CI) Total events: 0 (Other treatment), Heterogeneity: not applicable Test for overall effect: not applicab	. ,	0			Not estimable
24 Tar Subtotal (95% CI) Fotal events: 0 (Other treatment), Heterogeneity: not applicable Fest for overall effect: not applicab		0			Not estimable
25 Tazarotene					
		-	I -0.5 0 0.5 I		
		Favours oth	her treatment Favours placeb	0	(Continued

Study or subgroup	Other treatment n/N	Placebo n/N	Risk Difference H,Random,95% Cl	Weight	(Continued) Risk Difference H,Random,95% Cl
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatr	nent), 0 (Placebo)				
Heterogeneity: not applicabl	e				
Test for overall effect: not ap	plicable				
26 Theophylline 1% ointmer	it, twice daily				
Papakostantinou 2005	0/11	0/11		100.0 %	0.0 [-0.16, 0.16]
Subtotal (95% CI)	11	11	+	100.0 %	0.0 [-0.16, 0.16]
Total events: 0 (Other treatr	nent), 0 (Placebo)				
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 0$.	0 (P = 1.0)				
			-I -0.5 0 0.5 I		
		Favours ot	her treatment Favours placebo)	

Analysis 6.10. Comparison 6 Other treatment versus placebo, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 6 Other treatment versus placebo

Outcome: 10 Adverse events (systemic)

Study or subgroup Oth	ner treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Aloe vera extract 0.5% hydrophilic c	ream, three times p	er day			
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatment), 0 (F	Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applicable					
2 Anti-IL-8 monoclonal antibody crean	n				Not estimable
Subtotal (95% CI)	0	0			
Total events: 0 (Other treatment), 0 (F	Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applicable					
3 Betamethasone 17-valerate 21-aceta	te plus tretinoin plu	s salicylic acid			
Santoianni 2001	0/44	0/41	-	100.0 %	0.0 [-0.04, 0.04]
Subtotal (95% CI)	44	41	+	100.0 %	0.0 [-0.04, 0.04]
Total events: 0 (Other treatment), 0 (F	Placebo)				
Heterogeneity: not applicable	,				
Test for overall effect: $Z = 0.0$ (P = 1.0))				
4 Caffeine (topical) 10%, TD	,				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatment), 0 (F	Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applicable					
5 Calcipotriene 0.005% ointment + nic	cotinamide 0.05% oı	~ 0.1% or 0.7% or 1.4	ł%, twice daily		
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatment), 0 (F	Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applicable					
6 Dead Sea salts emollient lotion					Not estimable
Subtotal (95% CI)	0	0			
Total events: 0 (Other treatment), 0 (F	Placebo)				
Heterogeneity: not applicable					
Test for overall effect: not applicable					
7 Fish oil plus occlusion					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Other treatment), 0 (F	Placebo)				
			-I -0.5 0 0.5 I		
		Favours of	ther treatment Favours placebo	0	Continued

(Continued . . .)

Study or subgroup	Other treatment	Placebo	Risk Difference	Weight	Risk Difference
	n/N	n/N	M- H,Random,95% Cl		M: H,Random, C
Heterogeneity: not applicable Test for overall effect: not appli	cable				
B Herbal skin care (Dr Michael: Subtotal (95% CI) Fotal events: 0 (Other treatment Heterogeneity: not applicable Fest for overall effect: not appli P Hexafluoro-1,25-dihydroxyvit	0 nt), 0 (Placebo) cable	and skin conditioner), 0	twice daily		Not estimable
Durakovic 2001	0/15	0/15	=	100.0 %	0.0 [-0.12, 0.12]
Subtotal (95% CI) Total events: 0 (Other treatment Heterogeneity: not applicable Fest for overall effect: Z = 0.0 (, , ,	15	+	100.0 %	0.0 [-0.12, 0.12]
0 Indigo naturalis 1.4% ointme Lin 2007	ent 0/10	0/10	_	9.3 %	0.0 [-0.17, 0.17
Lin 2008	0/34	0/34	-	90.7 %	0.0 [-0.06, 0.06
Subtotal (95% CI) Total events: 0 (Other treatment Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 0.0 ($^{2} = 0.0, df = 1 (P = 1.00);$	44 I ² =0.0%	•	100.0 %	0.0 [-0.05, 0.05]
I Kukui nut oil, TD Subtotal (95% CI) Fotal events: 0 (Other treatmen Heterogeneity: not applicable Fest for overall effect: not appli	, , ,	0			Not estimable
2 <i>Mahonia aquifolium</i> (F Subtotal (95% CI) Total events: 0 (Other treatment Heterogeneity: not applicable Test for overall effect: not appli	0 nt), 0 (Placebo)	0			Not estimable
3 Methotrexate gel					
Sutton 2001	0/53	0/53	Ī	75.1 %	0.0 [-0.04, 0.04]
Syed 2001b	0/30	0/30	Ţ	24.9 %	0.0 [-0.06, 0.06]
Subtotal (95% CI) Total events: 0 (Other treatment	$^{2} = 0.0, df = 1 (P = 1.00);$	83 I ² =0.0%		100.0 %	0.0 [-0.03, 0.03]
Heterogeneity: $Tau^2 = 0.0$; Chi ² Test for overall effect: Z = 0.0 ((1 – 1.0)				
θ,					

Study or subgroup Other tre	atment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
Subtotal (95% CI) Total events: 0 (Other treatment), 0 (Placebo Heterogeneity: not applicable Test for overall effect: not applicable	0 >)	0			Not estimable
5 NG-monomethyl-L-arginine (L-NMMA) o Ormerod 2000	ream 0/17	0/17	-	100.0 %	0.0 [-0.11, 0.11]
Subtotal (95% CI) Total events: 0 (Other treatment), 0 (Placebo Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0)	17	17	•	100.0 %	0.0 [-0.11, 0.11]
6 Nicotinamide I.4%, twice daily Subtotal (95% CI) Total events: 0 (Other treatment), 0 (Placebo Heterogeneity: not applicable Test for overall effect: not applicable	0 D)	0			Not estimable
7 Oleum horwathiensis Lassus 1991	0/25	0/25	-	100.0 %	0.0 [-0.07, 0.07]
Subtotal (95% CI) Total events: 0 (Other treatment), 0 (Placebo Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0)	25	25	+	100.0 %	0.0 [-0.07, 0.07]
8 Omega-3-polyunsaturated fatty acids oint Subtotal (95% CI) Total events: 0 (Other treatment), 0 (Placebo Heterogeneity: not applicable Test for overall effect: not applicable	0	0			Not estimable
9 Platelet aggregation activating factor (PAF Wolska 1995)(Ro 24-0238) 0/52	0/52		100.0 %	0.0 [-0.04, 0.04]
Subtotal (95% CI) total events: 0 (Other treatment), 0 (Placebo Heterogeneity: not applicable test for overall effect: Z = 0.0 (P = 1.0)	52	52	•	100.0 %	0.0 [-0.04, 0.04]
0 Polymyxin B cream 200,000 U/g Subtotal (95% CI) Total events: 0 (Other treatment), 0 (Placebo Heterogeneity: not applicable Test for overall effect: not applicable	0	0			Not estimable
1 PTH (1-34) in Novasome A liposomal Holick 2003	cream, twice dai 0/15	ly 0/15	-	100.0 %	0.0 [-0.12, 0.12]

Study or subgroup	Other treatment	Placebo	Risk Difference M- H,Random,95%	Weight	(Continued) Risk Difference M- H,Random,95
	n/N	n/N	Ċl		Ċl
Subtotal (95% CI) Total events: 0 (Other treatmer Heterogeneity: not applicable Test for overall effect: Z = 0.0 (1		15	•	100.0 %	0.0 [-0.12, 0.12]
22 Sirolimus (topical) Subtotal (95% CI) Total events: 0 (Other treatmer Heterogeneity: not applicable Test for overall effect: not applic		0			Not estimable
23 Tacrolimus ointment Subtotal (95% CI) Total events: 0 (Other treatmer Heterogeneity: not applicable Test for overall effect: not applic		0			Not estimable
24 Tar Subtotal (95% CI) Total events: 0 (Other treatmer Heterogeneity: not applicable Test for overall effect: not applic		0			Not estimable
25 Tazarotene					
Krueger 1998	0/45	0/45	+	10.1 %	0.0 [-0.04, 0.04]
Weinstein 1996	0/216	0/108	•	89.9 %	0.0 [-0.01, 0.01]
Subtotal (95% CI) Total events: 0 (Other treatmer Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 0.0 (1	$P^2 = 0.0, df = 1 (P = 1.00);$	153 I ² =0.0%		100.0 %	0.0 [-0.01, 0.01]
26 Theophylline 1% ointment, t Subtotal (95% CI) Total events: 0 (Other treatmer Heterogeneity: not applicable Test for overall effect: not applic	0 nt), 0 (Placebo)	0			Not estimable

Analysis 7.1. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome I IAGI.

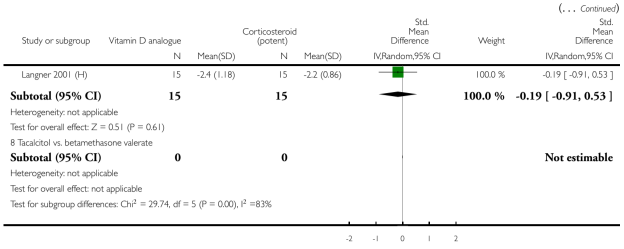
Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: I IAGI

Std Mean Difference	Weight	Std. Mean Difference	b	Corticosteroio (potent)	C	Vitamin D analogue	Study or subgroup
IV,Random,95% C		IV,Random,95% CI	Mean(SD)	Ν	Mean(SD)	Ν	
						thasone dipropionate	I Calcipotriol vs. betame
0.28 [-0.04, 0.60]	16.6 %		2.03 (1.06)	78	2.3 (0.81)	74	Fleming 2010 (H)
0.37 [0.24, 0.50]	45.3 %	-	1.9 (0.94)	476	2.24 (0.9)	480	Kaufmann 2002 (H)
0.56 [0.40, 0.72]	38.1 %	-	-3.44 (1.05)	312	-2.81 (1.21)	308	Papp 2003 (H)
0.43 [0.28, 0.58]	100.0 %	•		866		862	Subtotal (95% CI)
				50%	(P = 0.13); I ² =5	= 5.57 (P < 0.00001)	Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 2 Calcipotriol vs. betamet
-0.02 [-0.21, 0.17]	100.0 %		-2.39 (0.92)	207	-2.41 (0.94)	205	Molin 1997
-0.02 [-0.21, 0.17]	100.0 %	+		207		= 0.22 (P = 0.83)	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z =
Not estimable				0		netasone 0	3 Calcipotriol vs. desoxyn Subtotal (95% CI)
						able	Heterogeneity: not applic
							Test for overall effect: not
				100			4 Calcipotriol vs. diflorase
0.27 [0.02, 0.52]	100.0 %		-4.4 (.)	128	-4. (.)	128	Medansky 1996
-0.58 [-0.99, -0.18	100.0 %	◆	-3.3 (1.2)	128	-4.04 (1.31)	= 2.15 (P = 0.032)	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z = 5 Calcipotriol vs. fluocino Bruce 1994
-			-5.5 (1.2)		-1.01 (1.51)		
- 0.58 [-0.99, -0.18]	100.0 %		-3.55 (1.21)	47	-3.31 (1.12)	= 2.84 (P = 0.0046)	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z = 6 Calcitriol vs. betametha Camarasa 2003
L			5.55 (1.21)		5.51 (1.12)		
0.21 [-0.04, 0.45]	100.0 %			130		= 1.64 (P = 0.10)	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z = 7 Calcitriol vs. betametha

(Continued . . .)



Fav vitamin D analogue Fav corticosteroid (poten

Analysis 7.2. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 2 TSS

Study or subgroup	Vitamin D analogue		Corticosteroi (potent)	d	Diff	Std. Mean ference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Rando	om,95% Cl		IV,Random,95% CI
I Calcipotriol vs. betame	ethasone dipropionate							
Subtotal (95% CI)) 0		0					Not estimable
Heterogeneity: not appli	cable							
Test for overall effect: no	ot applicable							
2 Calcipotriol vs. betame	ethasone valerate							
Kragballe 1991a	342	2.31 (1.95)	342	2.82 (1.95)			100.0 %	-0.26 [-0.41, -0.11]
Subtotal (95% CI)	342		342		•		100.0 %	-0.26 [-0.41, -0.11]
Heterogeneity: not appli	cable							
Test for overall effect: Z	= 3.40 (P = 0.00067)							
3 Calcipotriol vs. desoxy	metasone							
Subtotal (95% CI)) 0		0					Not estimable
							1	
				-	2 -I C)	2	
				Favours vitami	n D analogue	Favours co	ticosteroid (poter	,
								(Continued)

Topical treatments for chronic plaque psoriasis (Review)

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N 128 128 16) 44 44 44	Mean(SD) 2.7 (1.5) 1.92 (1.48)	128 128	Mean(SD) 2.1 (1.5)	IV,Random,95% CI ➡ ←	100.0 % 100.0 %	Ⅳ,Random,95% C 0.40 [0.15, 0.65] 0.40 [0.15, 0.65]
128 (6) 44			2.1 (1.5)	•		-
128 (6) 44			2.1 (1.5)	•		
128 (6) 44			2.1 (1.5)	•		-
128 (6) 44			2.1 (1.5)	■		
16) 44	1.92 (1.48)	128		•	100.0 %	0.40 [0.15, 0.65
44	1.92 (1.48)					
44	1.92 (1.48)					
	1.92 (1.48)					
	1.92 (1.48)					
44		45	2.66 (1.48)		100.0 %	-0.50 [-0.92, -0.07]
		45		•	100.0 %	-0.50 [-0.92, -0.07]
I)						
te						
128	1.58 (0.82)	130	1.36 (0.82)		100.0 %	0.27 [0.02, 0.5
128		130		•	100.0 %	0.27 [0.02, 0.51
		100			10000 /0	0.2/[0.02, 0.51]
2)						
/						
0		0				Not estimable
63	3.06 (1.95)	63	2.3 (1.95)		85.5 %	0.39 [0.03, 0.74
11	2.77 (1.48)	П	1.92 (1.43)		14.5 %	0.56 [-0.29, 1.42
74		74		•	100.0 %	0.41 [0.09, 0.74
f = 1 ($P = 0.71$); $I^2 = 0.09$	%				
3)						
7, df =	= 4 (P = 0.00), I ² =	-89%				
	63 74 = (63 3.06 (1.95) 11 2.77 (1.48) 74 74 75 74 74 75 76 76 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77 77	63 3.06 (1.95) 63 11 2.77 (1.48) 11 74 74 74 5 = 1 (P = 0.71); 1 ² =0.0%	63 3.06 (1.95) 63 2.3 (1.95) 11 2.77 (1.48) 11 1.92 (1.43) 74 74 $= 1 (P = 0.71); 1^2 = 0.0\%$ $= 1, (P = 0.00), 1^2 = 89\%$	$63 3.06 (1.95) \qquad 63 2.3 (1.95) \\ 11 2.77 (1.48) \qquad 11 1.92 (1.43) \\ 74 \qquad 74 \qquad$	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

Favours vitamin D analogue

Favours corticosteroid (potent)

Analysis 7.3. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 3 PASI

I0.00 % 0.12 [-0.20, 0.44] 45.4 % 0.36 [0.23, 0.49] 38.2 % 0.47 [0.31, 0.63] 100.0 % 0.36 [0.22, 0.51] 26.7 % -0.03 [-0.23, 0.17] 41.9 % -0.18 [-0.33, -0.02] 27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % 0.15 [-0.73, 1.02]	IV,Random,95% CI	Mean(SD) 3.9 (3.8) -0.57 (0.3) -0.63 (0.27)	N 78 476 312 866 0%	Mean(SD) 4.3 (2.8) -0.46 (0.31) -0.49 (0.32)	74 480 308	I Calcipotriol vs. betame Fleming 2010 (H) Kaufmann 2002 (H)
45.4 % 0.36 [0.23, 0.49] 38.2 % 0.47 [0.31, 0.63] 100.0 % 0.36 [0.22, 0.51] 26.7 % -0.03 [-0.23, 0.17] 41.9 % -0.18 [-0.33, -0.02] 27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % 0.15 [-0.73, 1.02]		-0.57 (0.3)	476 312 866	-0.46 (0.31) -0.49 (0.32)	74 480 308	Fleming 2010 (H)
45.4 % 0.36 [0.23, 0.49] 38.2 % 0.47 [0.31, 0.63] 100.0 % 0.36 [0.22, 0.51] 26.7 % -0.03 [-0.23, 0.17] 41.9 % -0.18 [-0.33, -0.02] 27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % 0.15 [-0.73, 1.02]		-0.57 (0.3)	476 312 866	-0.46 (0.31) -0.49 (0.32)	480 308	
38.2 % 0.47 [0.31, 0.63] 100.0 % 0.36 [0.22, 0.51] 26.7 % -0.03 [-0.23, 0.17] 41.9 % -0.18 [-0.33, -0.02] 27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % 0.15 [-0.73, 1.02]		. ,	312 866	-0.49 (0.32)	308	Kaufmann 2002 (H)
100.0 % 0.36 [0.22, 0.51] 26.7 % -0.03 [-0.23, 0.17] 41.9 % -0.18 [-0.33, -0.02] 27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % -0.12 [-0.22, -0.02]	•	-0.63 (0.27)	866			14441114111 2002 (11)
26.7 % -0.03 [-0.23, 0.17] 41.9 % -0.18 [-0.33, -0.02] 27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % -0.12 [-0.22, -0.02] 100.0 % 0.15 [-0.73, 1.02]	•					Papp 2003 (H)
41.9 % -0.18 [-0.33, -0.02] 27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % -0.12 [-0.22, -0.02] 100.0 % 0.15 [-0.73, 1.02]	+		0%		862	Subtotal (95% CI)
41.9 % -0.18 [-0.33, -0.02] 27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % -0.12 [-0.22, -0.02] 100.0 % 0.15 [-0.73, 1.02]	+			$(P = 0.14); ^2 = 50$	= 4.81 (P < 0.00001)	Heterogeneity: Tau ² = 0.1 Test for overall effect: Z = 2 Calcipotriol vs. betame
27.4 % -0.11 [-0.30, 0.08] 4.0 % -0.16 [-0.66, 0.35] 100.0 % -0.12 [-0.22, -0.02]		-5.32 (6.02)	200	-5.5 (5.84)	201	Cunliffe 1992
4.0 % -0.16 [-0.66, 0.35] 100.0 % -0.12 [-0.22, -0.02] 100.0 % 0.15 [-0.73, 1.02]		3.06 (3.38)	316	2.5 (2.86)	316	Kragballe 1991a
100.0 % -0.12 [-0.22, -0.02]	-	3.5 (4.3)	207	3.1 (2.8)	205	Molin 1997
100.0 % 0.15 [-0.73, 1.02]		1.54 (3.61)	28	0.97 (3.61)	32	Vladimirov 1994
100.0 % 0.15 [-0.73, 1.02	•		751		754	Subtotal (95% CI)
L · · -				$P = 0.71$; $I^2 = 0.0$		Heterogeneity: $Tau^2 = 0.1$
L .					= 2.31 (P = 0.021)	Test for overall effect: Z =
L .					metasone	3 Calcipotriol vs. desoxyr
100.0 % 0.15 [-0.73, 1.02]		3.4 (1.93)	10	3.69 (1.9)	10	Kim 1994
			10		10	Subtotal (95% CI)
					cable	Heterogeneity: not applic
					= 0.32 (P = 0.75)	Test for overall effect: Z =
					one diacetate	4 Calcipotriol vs. diflorase
Not estimable			0		0	Subtotal (95% CI)
					cable	Heterogeneity: not applic
					t applicable	Test for overall effect: not
					onide	5 Calcipotriol vs. fluocino
Not estimable			0		0	Subtotal (95% CI)
					cable	Heterogeneity: not applic
					t applicable	Test for overall effect: not
					asone dipropionate	6 Calcitriol vs. betametha
100.0 % 0.39 [0.14, 0.63]		3.67 (3.79)	130	5.4 (5.06)	128	Camarasa 2003
100.0 % 0.39 [0.14, 0.63]	-		130		128	Subtotal (95% CI)

(Continued . . .)

Study or subgroup	Vitamin D analogue		ticosteroid (potent)		Dif	Std. Mean ference	Weight	(Continued Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Rando	om,95% Cl		IV,Random,95% CI
Heterogeneity: not applic	able							
Test for overall effect: Z =	= 3.07 (P = 0.0021)							
7 Calcitriol vs. betametha	isone valerate							
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applic	able							
Test for overall effect: not	t applicable							
8 Tacalcitol vs. betametha	asone valerate							
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applic	able							
Test for overall effect: not	t applicable							
Test for subgroup differer	nces: Chi ² = 34.88, df =	3 (P = 0.00), I ² = 9	91%					
				-2	- () 2		

Favours vitamin D analogue Favours corticosteroid (potent)

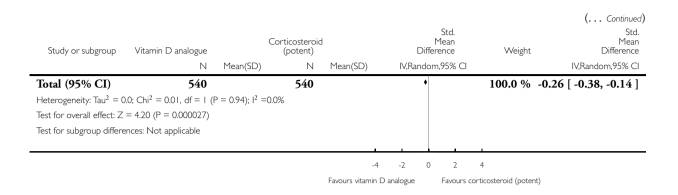
Analysis 7.4. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 4 PAGI

Study or subgroup	Vitamin D analogue		rticosteroio (potent)		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
I Calcipotriol vs. betame							
Subtotal (95% CI)			0				Not estimable
Heterogeneity: not applic							
Test for overall effect: no							
2 Calcipotriol vs. betame					_		
Cunliffe 1992	198	-2.52 (0.9)	198	-2.27 (1)	-	36.7 %	-0.26 [-0.46, -0.06]
Kragballe 1991a	342	-2.94 (0.64)	342	-2.77 (0.7)		63.3 %	-0.25 [-0.40, -0.10]
Subtotal (95% CI)	540		540		•	100.0 %	-0.26 [-0.38, -0.14]
Heterogeneity: $Tau^2 = 0$.	.0; $Chi^2 = 0.01$, $df = 1$ ($P = 0.94$); $I^2 = 0.0\%$	6				
Test for overall effect: Z =	= 4.20 (P = 0.000027)						
3 Calcipotriol vs. desoxyr	metasone						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	cable						
Test for overall effect: no	t applicable						
4 Calcipotriol vs. difloras	one diacetate						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	cable						
Test for overall effect: no	t applicable						
5 Calcipotriol vs. fluocino	onide						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	cable						
Test for overall effect: no	t applicable						
6 Calcitriol vs. betametha	asone dipropionate						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	cable						
Test for overall effect: no	t applicable						
7 Calcitriol vs. betametha	asone valerate						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	cable						
Test for overall effect: no	t applicable						
8 Tacalcitol vs. betametha	asone valerate						
Subtotal (95% CI)			0				Not estimable
Heterogeneity: not applie							
Test for overall effect: no	t applicable						
				-4	-2 0 2	4	
				Favours vitamin E	Danalogue Favours c	orticosteroid (poter	,
							(Continued



Analysis 7.5. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D analogue		Corticosteroi (potent)	d		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Ra	ndom,95% Cl		IV,Random,95% CI
l Calcipotriol vs. betame	thasone dipropionate							
Fleming 2010 (H)	74	2.3 (0.81)	78	2.03 (1.06)		-	16.6 %	0.28 [-0.04, 0.60]
Kaufmann 2002 (H)	480	2.24 (0.9)	476	1.9 (0.94)		-	45.3 %	0.37 [0.24, 0.50]
Papp 2003 (H)	308	-2.81 (1.21)	312	-3.44 (1.05)		•	38.1 %	0.56 [0.40, 0.72]
Subtotal (95% CI)	862		866			•	100.0 %	0.43 [0.28, 0.58]
Heterogeneity: $Tau^2 = 0$.01; Chi ² = 4.03, df = 2	(P = 0.13); I ² =	=50%					
Test for overall effect: Z	= 5.57 (P < 0.00001)							
2 Calcipotriol vs. betame	thasone valerate							
Cunliffe 1992	201	-5.5 (5.84)	200	-5.32 (6.02)		•	28.2 %	-0.03 [-0.23, 0.17]
Kragballe 1991a	342	2.31 (1.95)	342	2.82 (1.95)			36.4 %	-0.26 [-0.41, -0.11]
Molin 1997	205	-2.41 (0.94)	207	-2.39 (0.92)		•	28.6 %	-0.02 [-0.21, 0.17]
Vladimirov 1994	32	0.97 (3.61)	28	1.54 (3.61)		-	6.8 %	-0.16 [-0.66, 0.35]
					1 -2	0 2	4	
				 Favours vitamin		0 2 Eavours.co	4 orticosteroid (potent)	
						i avoui s co	acosteroid (poterit)	

(Continued . . .)

		Corticosteroid			Std. Mean	(Continued) Std. Mean	
Study or subgroup Vitamir	n D analogue N	Mean(SD)	(potent) N	Mean(SD)	Difference IV,Random,95% Cl	Weight	Difference IV,Random,95% CI
Subtotal (95% CI)	780		777		•	100.0 %	-0.12 [-0.26, 0.02]
Heterogeneity: $Tau^2 = 0.01$; Chi ²		$(P = 0.16); ^2 = 42$					
Test for overall effect: $Z = 1.68$ (P							
3 Calcipotriol vs. desoxymetasone							
Kim 1994	10	3.69 (1.9)	10	3.4 (1.93)	-	100.0 %	0.15 [-0.73, 1.02]
Subtotal (95% CI)	10		10		-	100.0 %	0.15 [-0.73, 1.02]
Heterogeneity: not applicable							
Test for overall effect: $Z = 0.32$ (P	= 0.75)						
4 Calcipotriol vs. diflorasone diace	tate						
Medansky 1996	128	-4. (.)	128	-4.4 (.)		100.0 %	0.27 [0.02, 0.52]
Subtotal (95% CI)	128		128		•	100.0 %	0.27 [0.02, 0.52]
Heterogeneity: not applicable							
Test for overall effect: $Z = 2.15$ (P	= 0.032)						
5 Calcipotriol vs. fluocinonide							
Bruce 1994	52	-4.04 (1.31)	47	-3.3 (1.2)		100.0 %	-0.58 [-0.99, -0.18]
Subtotal (95% CI)	52		47		•	100.0 %	-0.58 [-0.99, -0.18]
Heterogeneity: not applicable							
Test for overall effect: $Z = 2.84$ (P	= 0.0046)						
6 Calcitriol vs. betamethasone dip	ropionate						
Camarasa 2003	128	-3.31 (1.12)	130	-3.55 (1.21)		100.0 %	0.21 [-0.04, 0.45]
Subtotal (95% CI)	128		130		•	100.0 %	0.21 [-0.04, 0.45]
Heterogeneity: not applicable							
Test for overall effect: $Z = 1.64$ (P	= 0.10)						
7 Calcitriol vs. betamethasone vale							
Langner 2001 (H)	15	-2.4 (1.18)	15	-2.2 (0.86)		100.0 %	-0.19 [-0.91, 0.53]
Subtotal (95% CI)	15		15		•	100.0 %	-0.19 [-0.91, 0.53]
Heterogeneity: not applicable							
Test for overall effect: $Z = 0.51$ (P	,						
8 Tacalcitol vs. betamethasone val	erate						
Scarpa 1996	63	3.06 (1.95)	63	2.3 (1.95)	-	85.5 %	0.39 [0.03, 0.74]
Seidenari 1997 (H)	11	2.77 (1.48)	11	1.92 (1.43)		14.5 %	0.56 [-0.29, 1.42]
Subtotal (95% CI)	74		74		•	100.0 %	0.41 [0.09, 0.74]
Heterogeneity: Tau ² = 0.0; Chi ² =	0.14, df = 1 ($P = 0.7 I$); $I^2 = 0.0$	%				
Test for overall effect: $Z = 2.48$ (P	= 0.013)						
Test for subgroup differences: Chi	² = 44.40, df =	7 (P = 0.00), I ² =	=84%				
						L	
				-4	-2 0 2	4	

Analysis 7.6. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D analogue	Corticosteroid (potent)	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
I Calcipotriol vs. betamethasor	ne dipropionate				
Fleming 2010 (H)	6/79	5/83		3.6 %	0.02 [-0.06, 0.09]
Kaufmann 2002 (H)	39/480	22/476	-	22.6 %	0.04 [0.00, 0.07]
Papp 2003 (H)	27/308	17/313	-	13.2 %	0.03 [-0.01, 0.07]
Subtotal (95% CI)	867	872	•	39.4 %	0.03 [0.01, 0.06]
Total events: 72 (Vitamin D ana Heterogeneity: Tau ² = 0.0; Chi Test for overall effect: Z = 2.74 2 Calcipotriol vs. betamethason	$i^2 = 0.21, df = 2 (P = 0.90)$ 4 (P = 0.0061)	4 <i>//</i>			
Cunliffe 1992	21/205	17/204		6.8 %	0.02 [-0.04, 0.08]
Kragballe 1991a	15/345	15/345	+	23.3 %	0.0 [-0.03, 0.03]
Molin 1997	14/210	7/211	-	12.5 %	0.03 [-0.01, 0.07]
Subtotal (95% CI)	760	760	•	42.6 %	0.01 [-0.01, 0.04]
Test for overall effect: Z = 1.12 3 Calcipotriol vs. desoxymetase Subtotal (95% CI) Total events: 0 (Vitamin D anal Heterogeneity: not applicable Test for overall effect: not appli	one 0 logue), 0 (Corticosteroid (j	0 potent))			Not estimable
4 Calcipotriol vs. diflorasone di Medansky 1996	iacetate 6/134	6/134	—	8.8 %	0.0 [-0.05, 0.05]
,					
Subtotal (95% CI) Total events: 6 (Vitamin D anal Heterogeneity: not applicable Test for overall effect: Z = 0.0 5 Calcipotriol vs. fluocinonide		134 potent))	•	8.8 %	0.0 [-0.05, 0.05]
Subtotal (95% CI) Total events: 0 (Vitamin D anal Heterogeneity: not applicable	0 logue), 0 (Corticosteroid (j	0 potent))			Not estimable
		-0. Favours vitamin		osteroid (potent)	(Continued)

					(Continued
Study or subgroup	Vitamin D analogue	Corticosteroid (potent)	Risk Difference	Weight	Risk Difference
			M- H,Random,95%		M- H,Random,9
	n/N	n/N	Cl		Cl
Test for overall effect: not ap	•				
6 Calcitriol vs. betamethasor					
Camarasa 2003	6/128	9/130	-	6.6 %	-0.02 [-0.08, 0.03]
Subtotal (95% CI)	128	130	•	6.6 %	-0.02 [-0.08, 0.03]
Total events: 6 (Vitamin D a	nalogue), 9 (Corticosteroid (potent))			
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 0$.77 (P = 0.44)				
7 Calcitriol vs. betamethasor	ne valerate				
Langner 2001 (H)	1/15	0/15		0.8 %	0.07 [-0.10, 0.23]
Subtotal (95% CI)	15	15	-	0.8 %	0.07 [-0.10, 0.23]
Total events: I (Vitamin D a	nalogue), 0 (Corticosteroid (potent))			
Heterogeneity: not applicabl	le	<i>,,</i>			
Test for overall effect: $Z = 0$.79 (P = 0.43)				
8 Tacalcitol vs. betamethaso	ne valerate				
Scarpa 1996	13/76	13/76		1.5 %	0.0 [-0.12, 0.12]
Seidenari 1997 (H)	3/14	3/14		0.2 %	0.0 [-0.30, 0.30]
Subtotal (95% CI)	90	90	+	1.7 %	0.0 [-0.11, 0.11]
Total events: 16 (Vitamin D	analogue), 16 (Corticosteroid	d (potent))			
Heterogeneity: $Tau^2 = 0.0$; (Chi ² = 0.0, df = 1 (P = 1.00);	$ ^2 = 0.0\%$			
Test for overall effect: $Z = 0$.0 (P = 1.0)				
Total (95% CI)	1994	2001	•	100.0 %	0.02 [0.00, 0.03]
Total events: 151 (Vitamin D) analogue), 114 (Corticoster	oid (potent))			
Heterogeneity: $Tau^2 = 0.0$; ($Chi^2 = 6.50, df = 10 (P = 0.7)$	7); I ² =0.0%			
Test for overall effect: $Z = 2$.33 (P = 0.020)				
Test for subgroup difference	s: $Chi^2 = 4.58$, $df = 5$ (P = 0.	47), l ² =0.0%			
		-0.	5 -0.25 0 0.25 0.5		
		Favours vitamin	D analogue Favours cortic	osteroid (potent)	

Analysis 7.7. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D analogue	Corticosteroid (potent)	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Calcipotriol vs. betamethasone	dipropionate				
Kaufmann 2002 (H)	15/480	5/476	-	15.2 %	0.02 [0.00, 0.04]
Subtotal (95% CI)	480	476	•	15.2 %	0.02 [0.00, 0.04]
Total events: 15 (Vitamin D analo	gue), 5 (Corticosteroid	(potent))			
Heterogeneity: not applicable	5 /· (· · · ·			
Test for overall effect: Z = 2.25 (F	^o = 0.024)				
2 Calcipotriol vs. betamethasone	,				
Cunliffe 1992	4/205	2/204	+	9.2 %	0.01 [-0.01, 0.03]
Kragballe 1991a	2/345	1/345	•	51.5 %	0.00 [-0.01, 0.01]
Molin 1997	6/210	3/211	-	6.5 %	0.01 [-0.01, 0.04]
Subtotal (95% CI)	760	760		67.2 %	0.00 [0.00, 0.01]
Total events: 12 (Vitamin D analo Heterogeneity: Tau ² = 0.0; Chi ² =	0 / (4 <i>//</i>			
Test for overall effect: $Z = 1.13$ (F	^o = 0.26)				
3 Calcipotriol vs. desoxymetason	e				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D analog	ue), 0 (Corticosteroid (p	potent))			
Heterogeneity: not applicable					
Test for overall effect: not applica	ble				
4 Calcipotriol vs. diflorasone diac	etate				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D analog	ue), 0 (Corticosteroid (μ	potent))			
Heterogeneity: not applicable					
Test for overall effect: not applica	ble				
5 Calcipotriol vs. fluocinonide					
Bruce 1994	0/57	1/57		2.2 %	-0.02 [-0.06, 0.03]
Subtotal (95% CI)	57	57	•	2.2 %	-0.02 [-0.06, 0.03]
Total events: 0 (Vitamin D analog	ue), Ι (Corticosteroid (μ	potent))			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.73$ (F	P = 0.47)				
6 Calcitriol vs. betamethasone dip	propionate				
Camarasa 2003	2/128	1/130	ŧ	7.2 %	0.01 [-0.02, 0.03]
		-0	.5 -0.25 0 0.25 0).5	
		-u Favours vitamir		1.5 ticosteroid (potent)	
		ravours vitamin	i Dianaiogue Favours com	ucostei ola (potent)	(Continued

Study or subgroup	Vitamin D analogue	Corticosteroid (potent)	Risk Difference	Weight	Risk Difference
, , ,	n/N	n/N	M- H,Random,95% Cl	5	M- H,Random,959 Cl
Subtotal (95% CI)	128	130	•	7.2 %	0.01 [-0.02, 0.03]
Total events: 2 (Vitamin D a	nalogue), I (Corticosteroid (j	potent))			
Heterogeneity: not applicabl	le				
Test for overall effect: $Z = 0$	0.59 (P = 0.55)				
7 Calcitriol vs. betamethason	ne valerate				
Langner 2001 (H)	1/15	0/15	·	0.2 %	0.07 [-0.10, 0.23]
Subtotal (95% CI)	15	15		0.2 %	0.07 [-0.10, 0.23]
Total events: I (Vitamin D a	nalogue), 0 (Corticosteroid (j	potent))			
Heterogeneity: not applicabl	le				
Test for overall effect: $Z = 0$	0.79 (P = 0.43)				
8 Tacalcitol vs. betamethaso	ne valerate				
Scarpa 1996	0/76	0/76	+	7.7 %	0.0 [-0.03, 0.03]
Seidenari 1997 (H)	0/14	0/14		0.3 %	0.0 [-0.13, 0.13]
Subtotal (95% CI)	90	90	•	8.0 %	0.0 [-0.02, 0.02]
Total events: 0 (Vitamin D a	nalogue), 0 (Corticosteroid (j	potent))			
Heterogeneity: $Tau^2 = 0.0$; (Chi ² = 0.0, df = 1 (P = 1.00);	$ ^2 = 0.0\%$			
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
Total (95% CI)	1530	1528		100.0 %	0.01 [0.00, 0.01]
Total events: 30 (Vitamin D	analogue), 13 (Corticosteroid	l (potent))			
Heterogeneity: $Tau^2 = 0.0$; ($Chi^2 = 6.49, df = 8 (P = 0.59)$; I ² =0.0%			
Test for overall effect: $Z = I$.89 (P = 0.059)				
Test for subgroup difference	s: $Chi^2 = 4.28$, $df = 5$ (P = 0.	5 I), I ² =0.0%			
		-0.	5 -0.25 0 0.25 0.5	i	

Favours vitamin D analogue Favours corticosteroid (potent)

Analysis 7.8. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Vitamin D analogue	Corticosteroid (potent)	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
I Calcipotriol vs. betamethasone	dipropionate				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D analog	gue), 0 (Corticosteroid (p	otent))			
Heterogeneity: not applicable					
Test for overall effect: not applica	ble				
2 Calcipotriol vs. betamethasone	valerate				
Cunliffe 1992	6/205	6/204	•	7.3 %	0.00 [-0.03, 0.03]
Kragballe 1991a	1/345	2/345		80.5 %	0.00 [-0.01, 0.01]
Subtotal (95% CI)	550	549		87.8 %	0.00 [-0.01, 0.01]
Total events: 7 (Vitamin D analog	gue), 8 (Corticosteroid (p	otent))			
Heterogeneity: Tau ² = 0.0; Chi ² :	= 0.05, df = 1 (P = 0.82)	$ ^2 = 0.0\%$			
Test for overall effect: Z = 0.56 (H	P = 0.58)				
3 Calcipotriol vs. desoxymetason	e				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D analog	gue), 0 (Corticosteroid (p	otent))			
Heterogeneity: not applicable					
Test for overall effect: not applica	ble				
4 Calcipotriol vs. diflorasone diac	etate				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D analog	gue), 0 (Corticosteroid (p	otent))			
Heterogeneity: not applicable					
Test for overall effect: not applica	ble				
5 Calcipotriol vs. fluocinonide					
Bruce 1994	0/57	0/56	•	6.7 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	57	56	•	6. 7 %	0.0 [-0.03, 0.03]
Total events: 0 (Vitamin D analog	gue), 0 (Corticosteroid (p	otent))			
Heterogeneity: not applicable	- 10				
Test for overall effect: $Z = 0.0$ (P	,				
6 Calcitriol vs. betamethasone dip Camarasa 2003	4/128	3/130	+	4.9 %	0.01 [-0.03, 0.05 -
Subtotal (95% CI)	128	130	•	4.9 %	0.01 [-0.03, 0.05]
Total events: 4 (Vitamin D analog				202 /0	
,		.,			
		-	I -0.5 0 0.5 I		
		Favours vitamin	D analogue Favours cortico	osteroid (potent)	
					(Continued

Study or subgroup	Vitamin D analogue n/N	Corticosteroid (potent) n/N	Risk Difference H,Random,95% Cl	Weight	(Continued) Risk Difference H,Random,95% C
Heterogeneity: not applicab		10/1 N	Ci		Ci
Test for overall effect: $Z = 0$					
7 Calcitriol vs. betamethaso	· · · ·				
Langner 2001 (H)	0/15	0/15	+	0.5 %	0.0 [-0.12, 0.12]
Subtotal (95% CI)	15	15	+	0.5 %	0.0 [-0.12, 0.12]
Total events: 0 (Vitamin D a	nalogue), 0 (Corticosteroid (potent))			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
8 Tacalcitol vs. betamethaso	ne valerate				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D a	nalogue), 0 (Corticosteroid (ootent))			
Heterogeneity: not applicab	le				
Test for overall effect: not ap	oplicable				
Total (95% CI)	750	750		100.0 %	0.00 [-0.01, 0.01]
Total events: II (Vitamin D	analogue), II (Corticosteroi	d (potent))			
Heterogeneity: $Tau^2 = 0.0$; ($Chi^2 = 0.50, df = 4 (P = 0.97)$); I ² =0.0%			
Test for overall effect: $Z = 0$	0.43 (P = 0.67)				
Test for subgroup difference	s: $Chi^2 = 0.29$, $df = 3$ (P = 0.29)	96), I ² =0.0%			
		-	I -0.5 0 0.5 I		

Favours vitamin D analogue

Alogue Favours corticosteroid (potent)

Analysis 7.9. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D analogue	Corticosteroid (potent)	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
I Calcipotriol vs. betamethason	ne dipropionate				
Fleming 2010 (H)	8/79	7/83	+	9.4 %	0.02 [-0.07, 0.11]
Kaufmann 2002 (H)	54/480	23/476	-	13.3 %	0.06 [0.03, 0.10]
Papp 2003 (H)	53/308	27/313	-	12.1 %	0.09 [0.03, 0.14]
Subtotal (95% CI)	867	872	•	34.9 %	0.07 [0.04, 0.09]
Total events: 115 (Vitamin D an Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 4.72 2 Calcipotriol vs. betamethason	² = 1.73, df = 2 (P = 0.42 (P < 0.00001)	u <i>m</i>			
Cunliffe 1992	64/205	18/204	+	10.5 %	0.22 [0.15, 0.30]
Kragballe 1991a	37/345	35/345	÷	12.6 %	0.01 [-0.04, 0.05]
Molin 1997	49/207	21/210	+	10.8 %	0.14 [0.07, 0.21
Subtotal (95% CI)	757	759	•	33.9 %	0.12 [-0.02, 0.26
3 Calcipotriol vs. desoxymetasc Subtotal (95% CI) Total events: 0 (Vitamin D analo Heterogeneity: not applicable	0	0 potent))			Not estimable
Test for overall effect: not applie 4 Calcipotriol vs. diflorasone dia	acetate				
Subtotal (95% CI) Total events: 0 (Vitamin D analo Heterogeneity: not applicable Test for overall effect: not applic		0 potent))			Not estimable
5 Calcipotriol vs. fluocinonide Bruce 1994	10/57	4/56		7.4 %	0.10 [-0.02, 0.22
Subtotal (95% CI) Total events: 10 (Vitamin D ana Heterogeneity: not applicable	57 alogue), 4 (Corticosteroid	56 (potent))		7.4 %	0.10 [-0.02, 0.22]
		- I Favours vitamin	-0.5 0 0.5 I D analogue Favours corticc	osteroid (potent)	
					(Continued

Study or subgroup	Vitamin D analogue	Corticosteroid (potent)	Risk Difference M- H,Random,95%	Weight	Risk Difference M- H,Random,95
	n/N	n/N	Cl		Cl
Test for overall effect: $Z = I$.	70 (P = 0.088)				
6 Calcitriol vs. betamethasor	ne dipropionate				
Camarasa 2003	7/128	7/130	+	11.9 %	0.00 [-0.05, 0.06]
Subtotal (95% CI)	128	130	+	11.9 %	0.00 [-0.05, 0.06]
Total events: 7 (Vitamin D ar	nalogue), 7 (Corticosteroid (j	potent))			
Heterogeneity: not applicable	e				
Test for overall effect: $Z = 0$.	03 (P = 0.98)				
7 Calcitriol vs. betamethasor	ne valerate				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D ar	nalogue), 0 (Corticosteroid (p	potent))			
Heterogeneity: not applicable	e				
Test for overall effect: not ap	plicable				
8 Tacalcitol vs. betamethasor	ne valerate				
Scarpa 1996	2/76	3/76	1	11.8 %	-0.01 [-0.07, 0.04]
Subtotal (95% CI)	76	76	+	11.8 %	-0.01 [-0.07, 0.04]
Total events: 2 (Vitamin D ar	nalogue), 3 (Corticosteroid (j	potent))			
Heterogeneity: not applicable	e				
Test for overall effect: $Z = 0$.	46 (P = 0.65)				
Total (95% CI)	1885	1893	•	100.0 %	0.07 [0.02, 0.11]
Total events: 284 (Vitamin D	analogue), 145 (Corticoster	oid (potent))			
Heterogeneity: $Tau^2 = 0.00;$	Chi ² = 45.38, df = 8 (P<0.00	0001); I ² =82%			
Test for overall effect: $Z = 2$.	()				
Test for subgroup differences	s: $Chi^2 = 10.77$, $df = 4$ (P = 0	0.03), I ² =63%			

Favours vitamin D analogue Favours corticosteroid (potent)

Analysis 7.10. Comparison 7 Vitamin D analogues versus corticosteroid (potent), Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 7 Vitamin D analogues versus corticosteroid (potent)

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D analogue	Corticosteroid (potent)	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
Calcipotriol vs. betamethasc	one dipropionate				
Papp 2003 (H)	0/308	0/313	•	34.0 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	308	313		34.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: Z = 0.0 2 Calcipotriol vs. betamethasc) (P = 1.0)	potent))			
Cunliffe 1992	1/205	1/204	-	7.3 %	0.00 [-0.01, 0.01]
Kragballe 1991a	0/345	0/345	-	41.9 %	0.0 [-0.01, 0.01]
Molin 1997	0/207	0/210	-	15.4 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	757	759		64.7 %	0.00 [0.00, 0.00]
3 Calcipotriol vs. desoxymeta: Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app Calcipation vs. differences of	0 alogue), 0 (Corticosteroid (j plicable	0 potent))			Not estimable
4 Calcipotriol vs. diflorasone o	diacetate	0			
Subtotal (95% CI)					Not estimable
Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app	alogue), 0 (Corticosteroid (j plicable				Not estimable
Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app 5 Calcipotriol vs. fluocinonide Subtotal (95% CI)	alogue), 0 (Corticosteroid (j plicable				Not estimable Not estimable
Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app 5 Calcipotriol vs. fluocinonide Subtotal (95% CI) Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app	alogue), 0 (Corticosteroid (j plicable alogue), 0 (Corticosteroid (j plicable	potent)) 0			
Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not app	alogue), 0 (Corticosteroid (j plicable alogue), 0 (Corticosteroid (j plicable	potent)) 0		1.2 %	

(Continued ...)

Study or subgroup Vita	ımin D analogue	Corticosteroid (potent)	Risk Difference M-	Weight	Risk Difference M- H,Random,95% Cl	
	n/N	n/N	H,Random,95% Cl			
ubtotal (95% CI)	128	130	+	1.2 %	-0.01 [-0.04, 0.03]	
otal events: 2 (Vitamin D analogue)	, 3 (Corticosteroid (p	ootent))				
leterogeneity: not applicable						
est for overall effect: $Z = 0.43$ (P =	0.66)					
Calcitriol vs. betamethasone valera	te					
ubtotal (95% CI)	0	0			Not estimable	
otal events: 0 (Vitamin D analogue)	, 0 (Corticosteroid (p	ootent))				
leterogeneity: not applicable						
est for overall effect: not applicable						
Tacalcitol vs. betamethasone valera	te					
Scarpa 1996	7/76	7/76		0.2 %	0.0 [-0.09, 0.09]	
ubtotal (95% CI)	76	76	+	0.2 %	0.0 [-0.09, 0.09]	
otal events: 7 (Vitamin D analogue)	, 7 (Corticosteroid (p	ootent))				
leterogeneity: not applicable						
est for overall effect: $Z = 0.0$ (P = 1	.0)					
Cotal (95% CI)	1269	1278		100.0 %	0.00 [0.00, 0.00]	
otal events: 10 (Vitamin D analogue	e), II (Corticosteroid	(potent))				
leterogeneity: $Tau^2 = 0.0$; $Chi^2 = 0.0$	31, df = 5 (P = 1.00)	; l ² =0.0%				
est for overall effect: $Z = 0.05$ (P =	0.96)					
est for subgroup differences: Chi ² =	0.19, df = 3 (P = 0.9	98), I ² =0.0%				

-0.5 -0.25 0 0.25 0.5

Favours vitamin D analogue Favours corticosteroid (potent)

Analysis 8.1. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: I IAGI

Study or subgroup	Vitamin D analogue		Corticosteroid (v potent)					Std. Mean rence		Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Ra	Indon	n,95% C	1	IV,Random,95% CI
I Calcipotriol vs. Cloł Koo 2006	petasol propionate 21	3.16 (1.18)	21	2.93 (1.18)						0.19 [-0.42, 0.80]
				Favours vi	-10	-5	0	5 Favour	10	steroid (v potent)

Analysis 8.3. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: 3 PASI

Study or subgroup	Vitamin D analogue		Corticosteroid (v potent)					Std. Mean rence		Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,R	andor	n,95% Cl		IV,Random,95% CI
I Calcipotriol vs. Clot Landi 1993	betasol propionate 20	1.33 (1.4)	20	2.02 (2.6)	I		+	1	l	-0.32 [-0.95, 0.30]
				Favours vi	-10 tamin D	-5 analogue	0	5 Favours	10 corticost	teroid (v potent)

Analysis 8.4. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: 4 PAGI

Study or subgroup	Vitamin D analogue		Corticosteroid (v potent)			Std. Mean rence	Std. Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% Cl		IV,Random,95% CI	
l Calcipotriol vs. Clo Koo 2006	betasol propionate 21	3.29 (1.3)	21	2.74 (1.3)	-	I	0.42 [-0.20, 1.03]	
				Favours vit	-10 -5 0 amin D analogue	5 Favours co	10 prticosteroid (v potent)	

Analysis 8.5. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D analogue		Corticosteroid (v potent)			Diff	Std. Mean erence		Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		IV,Rando	m,95% Cl	CI	IV,Random,95% CI	
I Calcipotriol vs. Clo	betasol propionate									
Koo 2006	21	3.16 (1.18)	21	2.93 (1.18)					51.1 %	0.19 [-0.42, 0.80]
Landi 1993	20	1.33 (1.4)	20	2.02 (2.6)		-	I		48.9 %	-0.32 [-0.95, 0.30]
Total (95% CI)	41		41			•		1	00.0 %	-0.06 [-0.57, 0.44]
Heterogeneity: Tau ²	= 0.03; Chi ² = 1.35, d	f = I (P = 0.25)); I ² =26%							
Test for overall effect	Z = 0.24 (P = 0.81)									
Test for subgroup diff	erences: Not applicab	le								
					-10	-5 0	5	10		
				Favours vitar	nin D an	alogue	Favours	corticost	eroid (v pote	ent)

Analysis 8.6. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D analogue	Corticosteroid (v potent)	Risk Difference M-	Risk Difference M-	
	n/N	n/N	H,Random,95% Cl	H,Random,95% Cl	
I Calcipotriol vs. Clobetas	ol propionate				
Koo 2006	0/21	0/21	+	0.0 [-0.09, 0.09]	
			-I -0.5 0 0.5 I		
		Favou	rs vitamin Dianalogue Eavours conticost	eroid (v. potent)	

Favours vitamin D analogue Favours corticosteroid (v potent)

Analysis 8.7. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D analogue	Corticosteroid (v potent)	Risk Difference M-	Risk Difference M-
	n/N	n/N	H,Random,95% Cl	H,Random,95% Cl
l Calcipotriol vs. Clobetaso	ol propionate			
Коо 2006	0/21	0/21	+	0.0 [-0.09, 0.09]
				•
			-1 -0.5 0 0.5	I
		Favours	vitamin D analogue Favours cor	-ticosteroid (v potent)

Analysis 8.8. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Vitamin D analogue	Corticosteroid (v potent)	Risk Difference M	2 -	Risk Difference M-
	n/N	n/N	H,Random, C	,95%]	H,Random,95% Cl
I Calcipotriol vs. Clobetase	ol propionate				
Koo 2006	0/21	0/21	+		0.0 [-0.09, 0.09]
			-1 -0.5 0	0.5 I	
		Favour	rs vitamin D analogue Fa	avours corticosteroid (v pot	ent)

Analysis 8.9. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D analogue	Corticosteroid (v potent)	Risk Difference M-	Risk Difference M-
	n/N	n/N	H,Random,95% Cl	H,Random,95% Cl
I Calcipotriol vs. Clobetas	ol propionate			
Landi 1993	0/20	1/20		-0.05 [-0.18, 0.08]
				1
			-I -0.5 0 0.5	I
		Favours	vitamin D analogue Favours c	orticosteroid (v potent)

Analysis 8.10. Comparison 8 Vitamin D analogues versus corticosteroid (very potent), Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 8 Vitamin D analogues versus corticosteroid (very potent)

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D analogue	Corticosteroid (v potent)	Risk Difference M-	Risk Difference M-
	n/N	n/N	H,Random,95% Cl	H,Random,95% Cl
I Calcipotriol vs. Clobetasc Landi 1993	ol propionate 0/20	1/20	_	-0.05 [-0.18, 0.08]
			-1 -0.5 0 0.5 1	

Favours vitamin D analogue Favours corticosteroid (v potent)

Analysis 9.1. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: I IAGI

Study or subgroup	Vitamin D combined with corticosteroid N	Mean(SD)	Corticosteroid N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
Calcipotriol + betamethasc	one dipropionate v	. betamethasc	ne dipropionate				
Douglas 2002	369	-3.72 (0.96)	363	-3.25 (1.1)	-	38.7 %	-0.46 [-0.60, -0.31]
Fleming 2010 (H)	150	1.85 (1.05)	78	2.03 (1.06)		17.0 %	-0.17 [-0.44, 0.10]
Kaufmann 2002 (H)	490	1.5 (0.9)	476	1.9 (0.94)	-	44.3 %	-0.43 [-0.56, -0.31]
Subtotal (95% CI)	1009		917		•	100.0 % -0	.40 [-0.52, -0.27]
Heterogeneity: $Tau^2 = 0.01$; ($Chi^2 = 3.44, df = 2$	$(P = 0.18); I^2$	=42%				
Test for overall effect: $Z = 6.1$	2 (P < 0.00001)						
2 Calcipotriol + betamethasc	one dipropionate ve	. clobetasol pr	opionate				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable							
Test for overall effect: not app	olicable						
3 Calcipotriol + clobetasol pr	ropionate vs. clobe	asol propiona	te				
Koo 2006	44	2.11 (1.18)	21	2.93 (1.18)		100.0 %	-0.69 [-1.22, -0.15]
Subtotal (95% CI)	44		21		-	100.0 % -0	.69 [-1.22, -0.15]
Heterogeneity: not applicable							
Test for overall effect: $Z = 2.5$	52 (P = 0.012)						
Test for subgroup differences:	Chi ² = 1.07, df =	(P = 0.30),	2 =6%				
				-2	-1 0 1	2	

Favours vitamin D combination

Favours corticosteroid

Analysis 9.2. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 2 TSS

Study or subgroup	Vitamin D combined with corticosteroid	Co			Std. Mean Difference				Std. Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% C			5% CI	I IV,Random,95% CI	
Calcipotriol + betame	ethasone dipropionate vs	betamethasone diprop	ionate							
2 Calcipotriol + betame	ethasone dipropionate vs	clobetasol propionate								
Menter 2009	62	1.65 (0.91)	60	1.25 (0.86)				-		0.45 [0.09, 0.81]
3 Calcipotriol + clobeta	usol propionate vs. clobet	asol propionate								
·										
									<u> </u>	
					-2 -	.	0	I.	2	
				Favours vitamin	D combin	ation	Fa	avours	corticost	eroid

Analysis 9.3. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 3 PASI

Study or subgroup	Vitamin D combined with corticosteroid N	Mean(SD)	Corticosteroid N	Mean(SD)		Std. Mean rence 95% Cl	Weight	Std. Mean Difference IV.Random,95% CI
l Calcipotriol + betamethasc		× /		(JUL)	TV,T Candon	1,7570 Cl		
Douglas 2002	342	2.5 (2.5)	343	3.9 (3.7)	-		37.9 %	-0.44 [-0.59, -0.29]
Fleming 2010 (H)	147	3.1 (2.8)	78	3.9 (3.8)			4. %	-0.25 [-0.53, 0.02]
0 ()		~ /		. ,	_			
Kaufmann 2002 (H)	490	-0.71 (0.26)	476	-0.57 (0.3)	-		48.0 %	-0.50 [-0.63, -0.37]
Subtotal (95% CI)	979		89 7		•		100.0 % -0	.44 [-0.55, -0.33]
Heterogeneity: $Tau^2 = 0.00$; ($Chi^2 = 2.58, df = 2$	(P = 0.28); I ² =	=22%					
Test for overall effect: $Z = 7.9$	5 (P < 0.00001)							
2 Calcipotriol + betamethaso	ne dipropionate vs	. clobetasol pro	pionate					
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicable								
Test for overall effect: not app	licable							
3 Calcipotriol + clobetasol pr	opionate vs. clobe	asol propionate	e					
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicable								
Test for overall effect: not app	licable							
Total (95% CI)	979		89 7		•		100.0 % -0	.44 [-0.55, -0.33]
Heterogeneity: $Tau^2 = 0.00$; ($Chi^2 = 2.58, df = 2$	$(P = 0.28); I^2 =$	=22%					
Test for overall effect: $Z = 7.9$	5 (P < 0.00001)							
Test for subgroup differences:	Not applicable							
				-2	-1 0	I.	2	
			Fa	wours vitamin D c	ombination	Favours co	rticosteroid	

Analysis 9.4. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 4 PAGI

Study or subgroup	Vitamin D combined with corticosteroid	C	orticosteroid		Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% CI
Calcipotriol + betame	ethasone dipropionate vs.	betamethasone diproj	pionate			
2 Calcipotriol + betame	ethasone dipropionate vs.	clobetasol propionate				
3 Calcipotriol + clobeta	sol propionate vs. clobeta	sol propionate				
Koo 2006	44	2.37 (1.3)	21	2.74 (1.3)	+	-0.28 [-0.80, 0.24]
					-10 -5 0 5 10	
				Favours vitamin	D combination Favours corti	costeroid

Analysis 9.5. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D combined with corticosteroid N	Mean(SD)	Corticosteroid N	Mean(SD)	Std. Mean Difference IV,Random,95% CI	Weight	Std. Mean Difference IV,Random,95% CI
I Calcipotriol + betamethason	e dipropionate ve	s. betamethaso	ne dipropionate				
Douglas 2002	369	-3.72 (0.96)	363	-3.25 (1.1)	•	38.7 %	-0.46 [-0.60, -0.31]
Fleming 2010 (H)	150	1.85 (1.05)	78	2.03 (1.06)	-	17.0 %	-0.17 [-0.44, 0.10]
Kaufmann 2002 (H)	490	1.5 (0.9)	476	1.9 (0.94)	-	44.3 %	-0.43 [-0.56, -0.31]
Subtotal (95% CI)	1009		917		•	100.0 % -	0.40 [-0.52, -0.27]
Heterogeneity: Tau ² = 0.01; Cł	$mi^2 = 3.44$, df = 2	$(P = 0.18); I^2$	=42%				
Test for overall effect: Z = 6.12	(P < 0.00001)						
2 Calcipotriol + betamethason	e dipropionate ve	s. clobetasol pr	opionate				
Menter 2009	62	1.65 (0.91)	. 60	1.25 (0.86)	-	100.0 %	0.45 [0.09, 0.81]
Subtotal (95% CI)	62		60		•	100.0 %	0.45 [0.09, 0.81]
Heterogeneity: not applicable							
Test for overall effect: Z = 2.45	(P = 0.0 4)						
3 Calcipotriol + clobetasol pro	pionate vs. clobe	tasol propiona	te				
Коо 2006	44	2.11 (1.18)	21	2.93 (1.18)		100.0 %	-0.69 [-1.22, -0.15]
Subtotal (95% CI)	44		21		•	100.0 % -	0.69 [-1.22, -0.15]
Heterogeneity: not applicable							
Test for overall effect: Z = 2.52	(P = 0.0 2)						
		= 2 (P = 0.00)	$ ^2 = 90\%$				
Test for subgroup differences: C	∠ni~ — 20.80, at -	2 (1 0.00),					

Favours vitamin D combination Favours corticosteroid

Analysis 9.6. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D combined with corticosteroid	Corticosteroid	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
Calcipotriol + betamethason	e dipropionate vs. betame	thasone dipropionate			
Douglas 2002	28/372	21/365		19.3 %	0.02 [-0.02, 0.05]
Fleming 2010 (H)	13/162	5/83		5.7 %	0.02 [-0.05, 0.09]
Kaufmann 2002 (H)	13/490	22/476	-	44.7 %	-0.02 [-0.04, 0.00]
Subtotal (95% CI)	1024	924	+	69. 7 %	0.00 [-0.03, 0.03]
Total events: 54 (Vitamin D cor Heterogeneity: Tau ² = 0.00; Ch Test for overall effect: $Z = 0.01$ 2 Calcipotriol + betamethason	$hi^2 = 3.86$, df = 2 (P = 0.1 (P = 0.99)	5); I ² =48%			
Menter 2009	0/62	0/60	+	25.1 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	62	60	+	25.1 %	0.0 [-0.03, 0.03]
Total events: 0 (Vitamin D com Heterogeneity: not applicable Test for overall effect: $Z = 0.0$ (3 Calcipotriol + clobetasol prop	(P = 1.0)				
Коо 2006	0/44	0/21		5.2 %	0.0 [-0.07, 0.07]
Subtotal (95% CI)	44	21	-	5.2 %	0.0 [-0.07, 0.07]
Total events: 0 (Vitamin D com Heterogeneity: not applicable Test for overall effect: $Z = 0.0$ (), 0 (Corticosteroid)			
Total (95% CI)	1130	1005	+	100.0 %	0.00 [-0.02, 0.01]
Total events: 54 (Vitamin D cor Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 0.52	² = 3.87, df = 4 (P = 0.42	, , ,			

Favours vitamin D combination Favours corticosteroid

Analysis 9.7. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D combined with corticosteroid	Corticosteroid	Risk Difference M-	Risk Difference M-
	n/N	n/N	H,Random,95% Cl	H,Random,95% Cl
I Calcipotriol + betamethasone of	dipropionate vs. betamethasone	dipropionate		
Kaufmann 2002 (H)	3/490	5/476	+	0.00 [-0.02, 0.01]
2 Calcipotriol + betamethasone of	dipropionate vs. clobetasol prop	onate		
Menter 2009	0/62	0/60		0.0 [-0.03, 0.03]
3 Calcipotriol + clobetasol propie	onate vs. clobetasol propionate			
Koo 2006	0/44	0/21		0.0 [-0.07, 0.07]
			-0.2 -0.1 0 0.1 0.2	
		Favours vitan	nin D combination Favours corticoste	eroid

Analysis 9.8. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Vitamin D combined with corticosteroid	Corticosteroid		Risk Difference M-	Risk Difference M-
	n/N	n/N	ŀ	H,Random,95% Cl	H,Random,95% Cl
I Calcipotriol + betamethasc	one dipropionate vs. betamethasor	ne dipropionate			
2 Calcipotriol + betamethas	one dipropionate vs. clobetasol pro	opionate			
Menter 2009	0/62	0/60		+	0.0 [-0.03, 0.03]
3 Calcipotriol + clobetasol p	ropionate vs. clobetasol propionat	e			
Koo 2006	0/44	0/21		+	0.0 [-0.07, 0.07]
			-1 -0.5	0 0.5 I	
		Favours vitar	min D combinatio	n Favours corticos	steroid

Analysis 9.9. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D combined with corticosteroid	Corticosteroid	Risk Difference M- H,Random,95%	Weight	Risk Difference M- H,Random,95%
	n/N	n/N	Cl		Cl
Calcipotriol + betamethason		1 1		2 / 2 2/	
Douglas 2002	30/372	17/365		36.2 %	0.03 [0.00, 0.07]
Fleming 2010 (H)	12/160	7/83		8.5 %	-0.01 [-0.08, 0.06]
Kaufmann 2002 (H)	29/490	23/476	-	55.3 %	0.01 [-0.02, 0.04]
Subtotal (95% CI)	1022	924	•	100.0 %	0.02 [0.00, 0.04]
Total events: 71 (Vitamin D cor	mbined with corticostero	id), 47 (Corticosteroid)			
Heterogeneity: $Tau^2 = 0.0$; Chi	² = 1.59, df = 2 (P = 0.45	5); l ² =0.0%			
Test for overall effect: $Z = 1.63$	(P = 0.10)				
2 Calcipotriol + betamethason	e dipropionate vs. clobeta	asol propionate			
Menter 2009	4/62	6/60		100.0 %	-0.04 [-0.13, 0.06]
Subtotal (95% CI)	62	60		100.0 %	-0.04 [-0.13, 0.06]
Total events: 4 (Vitamin D com	bined with corticosteroic	l), 6 (Corticosteroid)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.71$	(P = 0.48)				
3 Calcipotriol + clobetasol pro	pionate vs. clobetasol pro	pionate			
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D com	bined with corticosteroic	l), 0 (Corticosteroid)			
Heterogeneity: not applicable					
Test for overall effect: not appli	cable				
Test for subgroup differences: C	$Chi^2 = 1.09, df = 1 (P = 0)$	0.30), I ² =8%			
		-0.	2 -0.1 0 0.1 0.2		
		Favours vitamin D	combination Favours cortic	osteroid	

Analysis 9.10. Comparison 9 Vitamin D combined with corticosteroid versus corticosteroid, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 9 Vitamin D combined with corticosteroid versus corticosteroid

Outcome: 10 Adverse events (systemic)

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-

Study or subgroup	Vitamin D combined with corticosteroid	Corticosteroid	Risk Difference M- H.Random,95%	Risk Difference M- H.Random,95%
	n/N	n/N	Cl	Cl
I Calcipotriol + betamethasor	ne dipropionate vs. betamethasor	ne dipropionate		
Douglas 2002	0/372	0/365	+	0.0 [-0.01, 0.01]
2 Calcipotriol + betamethasor	ne dipropionate vs. clobetasol pro	ppionate		
3 Calcipotriol + clobetasol pro	pionate vs. clobetasol propionat	e		
			-0.2 -0.1 0 0.1 0.2	
		Favours vita	min D combination Favours cortice	osteroid

Analysis 10.1. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: I IAGI

Std. Mean Difference IV,Random,95% CI	Weight	Std. Mean ifference dom,95% Cl		Mean(SD)	Dithranol N	Mean(SD)	Vitamin D analogue N	Study or subgroup
							bl	I Calcipotriol vs. dithranc
-0.64 [-0.83, -0.46]	27.1 %		-	-2.29 (0.97)	227	-2.81 (0.6)	231	Berth Jones 1992b
-0.52 [-0.83, -0.21]	24.8 %			-2.6 (1.52)	77	-3.4 (1.52)	89	Christensen 1999
0.46 [0.03, 0.89]	22.0 %			-3.6 (1.19)	40	-2.96 (1.53)	46	Van de Kerkhof 2006
-0.86 [-1.10, -0.61]	26.1 %			-1.69 (0.96)	131	-2.48 (0.88)	153	Wall 1998
-0.43 [-0.85, -0.01]	100.0 %	-	•	-2.44 (0.45)	475 l ² =89% 54	3 (P<0.00001); -2.19 (0.52)	l 6; Chi ² = 28.06, df = 3	Subtotal (95% CI) Heterogeneity: Tau ² = 0. Test for overall effect: Z = 2 Calcitriol vs. dithranol Hutchinson 2000
0.51 [0.13, 0.88]	100.0 %	•			54		able	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z = 3 Tacalcitol vs. dithranol
Not estimable					0 ² =91%	F I (P = 0.00), I	able applicable	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: not Test for subgroup differen

Analysis 10.2. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 2 TSS

Study or subgroup	Vitamin D analogue		Dithranol		Std. Mean Difference	Weight	Std. Mean Difference
study of subgroup	N	Mean(SD)	N	Mean(SD)	IV,Random,95% Cl	, reight	IV,Random,95% CI
I Calcipotriol vs. dithrano	I						
Christensen 1999	89	2.38 (1.65)	77	3.72 (1.65)	•	58.0 %	-0.81 [-1.13, -0.49]
Grattan 1997 (H)	22	1.8 (2)	22	2.2 (2.6)		42.0 %	-0.17 [-0.76, 0.42]
Subtotal (95% CI)	111		99		•	100.0 %	-0.54 [-1.16, 0.08]
Heterogeneity: $Tau^2 = 0.1$	5; Chi ² = 3.48, df = 1	$(P = 0.06); I^2 =$	71%				
Test for overall effect: Z =	: 1.71 (P = 0.087)						
2 Calcitriol vs. dithranol							
Hutchinson 2000	60	1.65 (0.76)	54	1.54 (0.95)	H	100.0 %	0.13 [-0.24, 0.50]
Subtotal (95% CI)	60		54		+	100.0 %	0.13 [-0.24, 0.50]
Heterogeneity: not applica	able						
Test for overall effect: Z =	: 0.68 (P = 0.50)						
3 Tacalcitol vs. dithranol							
Farkas 1999	42	-6.1 (2.4)	42	-5.7 (2.1)		100.0 %	-0.18 [-0.60, 0.25]
Subtotal (95% CI)	42		42		•	100.0 %	-0.18 [-0.60, 0.25]
Heterogeneity: not applica	able						
Test for overall effect: Z =	: 0.80 (P = 0.42)						
Test for subgroup differen	ces: Chi ² = 3.53, df = 2	$P = (P = 0.17), ^2$	=43%				
				ı		1	
				-4	-2 0 2	4	

Analysis 10.3. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 3 PASI

Study or subgroup	Vitamin D analogue		Dithranol		Std. Mean Difference	Std. Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% CI	
I Calcipotriol vs. dithranol							
Berth Jones 1992b	214	3.4 (2.7)	208	4.7 (4.4)	*	-0.36 [-0.55, -0.16]	
Monastirli 2000	35	2.58 (1.18)	35	0.34 (0.51)	+	2.44 [1.81, 3.06]	
Van de Kerkhof 2006	54	-0.56 (0.37)	52	-0.63 (0.3)	+	0.21 [-0.18, 0.59]	
2 Calcitriol vs. dithranol							
Hutchinson 2000	60	4.2 (3.9)	54	5.2 (5.5)	+	-0.21 [-0.58, 0.16]	
3 Tacalcitol vs. dithranol							
Farkas 1999	42	4.16 (3.22)	42	4.38 (3.05)	+	-0.07 [-0.50, 0.36]	
						1	
					-10 -5 0 5 1	0	

Analysis 10.4. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 4 PAGI

Study or subgroup	Vitamin D analogue		Dithranol		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
I Calcipotriol vs. dithranol							
Berth Jones 1992b	231	-2.72 (0.74)	227	-2.31 (1)		52.6 %	-0.47 [-0.65, -0.28]
Van de Kerkhof 2006	46	-2.89 (1.68)	40	-3.53 (1.43)	-	47.4 %	0.40 [-0.02, 0.83]
Subtotal (95% CI)	277		267		+	100.0 %	-0.05 [-0.90, 0.80]
Heterogeneity: $Tau^2 = 0.35$	5; Chi ² = 13.35, df = 1	(P = 0.00026);	l ² =93%				
Test for overall effect: Z =	0.12 (P = 0.90)						
2 Calcitriol vs. dithranol							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applical	ble						
Test for overall effect: not a	applicable						
3 Tacalcitol vs. dithranol							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applical	ble						
Test for overall effect: not a	applicable						
Test for subgroup difference	es: Not applicable						
				-4	-2 0 2	4	

Analysis 10.5. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D analogue		Dithranol		Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% CI
l Calcipotriol vs. dithranol						
Berth Jones 1992b	231	-2.81 (0.6)	227	-2.29 (0.97)		-0.64 [-0.83, -0.46]
Christensen 1999	89	-3.4 (1.52)	77	-2.6 (1.52)	- _	-0.52 [-0.83, -0.21]
Grattan 1997 (H)	22	1.8 (2)	22	2.2 (2.6)		-0.17 [-0.76, 0.42]
Monastirli 2000	35	2.58 (1.18)	35	0.34 (0.51)		→ 2.44 [I.81, 3.06]
Van de Kerkhof 2006	46	-2.96 (1.53)	40	-3.6 (1.19)		0.46 [0.03, 0.89]
Wall 1998	153	-2.48 (0.88)	131	-1.69 (0.96)		-0.86 [-1.10, -0.61]
2 Calcitriol vs. dithranol						
Hutchinson 2000	60	-2.19 (0.52)	54	-2.44 (0.45)		0.5 [0. 3, 0.88]
3 Tacalcitol vs. dithranol						
Farkas 1999	42	-6.1 (2.4)	42	-5.7 (2.1)		-0.18 [-0.60, 0.25]
					-2 -1 0 1	2

Analysis 10.6. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D analogue	Dithranol	Risk Difference M-	Weight	Risk Difference M
	n/N	n/N	H,Random,95% Cl		H,Random, C
l Calcipotriol vs. dithranol					
Christensen 1999	0/89	5/82	-	37.4 %	-0.06 [-0.12, -0.01
Grattan 1997 (H)	3/25	3/25		3.6 %	0.0 [-0.18, 0.18
Monastirli 2000	0/35	0/35	•	39.7 %	0.0 [-0.05, 0.05]
Van de Kerkhof 2006	17/54	11/52	+	4.2 %	0.10 [-0.06, 0.27]
Van der Vleuten 1995	0/10	0/10		3.8 %	0.0 [-0.17, 0.17
Subtotal (95% CI)	213	204	•	88.7 %	-0.01 [-0.07, 0.04]
Total events: 20 (Vitamin D a	analogue), 19 (Dithranol)				
Heterogeneity: $Tau^2 = 0.00;$	$Chi^2 = 6.25, df = 4 (P = 0.18)$	3); I ² =36%			
Test for overall effect: $Z = 0.4$	48 (P = 0.63)	,			
2 Calcitriol vs. dithranol					
Hutchinson 2000	12/60	16/54		4.6 %	-0.10 [-0.25, 0.06
Subtotal (95% CI)	60	54	•	4.6 %	-0.10 [-0.25, 0.06
Total events: 12 (Vitamin D a	analogue), 16 (Dithranol)				
Heterogeneity: not applicable	e				
Test for overall effect: $Z = I$.	19 (P = 0.23)				
3 Tacalcitol vs. dithranol					
Farkas 1999	4/42	5/42	-	6.6 %	-0.02 [-0.16, 0.11
Subtotal (95% CI)	42	42	+	6.6 %	-0.02 [-0.16, 0.11
Total events: 4 (Vitamin D an	nalogue), 5 (Dithranol)				
Heterogeneity: not applicable	e				
Test for overall effect: $Z = 0.2$	35 (P = 0.72)				
Total (95% CI)	315	300	•	100.0 %	-0.02 [-0.06, 0.01]
Total events: 36 (Vitamin D a	analogue), 40 (Dithranol)				
Heterogeneity: $Tau^2 = 0.0$; C	^c hi ² = 5.70, df = 6 (P = 0.46)	; l ² =0.0%			
Test for overall effect: $Z = 1$.	41 (P = 0.16)				
	$: Chi^2 = 0.95, df = 2 (P = 0.6)$	(2) 12 -0.000			

Analysis 10.7. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D analogue	Dithranol	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
I Calcipotriol vs. dithranol					
Berth Jones 1992b	4/239	12/239	-	33.0 %	-0.03 [-0.07, 0.00]
Lister 1997	2/89	6/82	-	14.7 %	-0.05 [-0.11, 0.01]
Monastirli 2000	0/35	0/35	+	18.7 %	0.0 [-0.05, 0.05]
Van de Kerkhof 2006	7/54	3/52	-	6.1 %	0.07 [-0.04, 0.18]
Van der Vleuten 1995	0/10	0/10	-	2.6 %	0.0 [-0.17, 0.17]
Wall 1998	9/161	20/145	-	13.9 %	-0.08 [-0.15, -0.02]
Subtotal (95% CI)	588	563	•	88.9 %	-0.03 [-0.06, 0.00]
Test for overall effect: Z = 1.69 2 Calcitriol vs. dithranol Hutchinson 2000	9 (P = 0.092) 1/60	4/54	-	. %	-0.06 [-0.13, 0.02]
Subtotal (95% CI)	60	54	•	11.1 %	-0.06 [-0.13, 0.02]
Total events: 1 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: Z = 1.46 3 Tacalcitol vs. dithranol	0, ()				
5 Tacalciloi VS. Ultriranoi	0	0			Not estimable
Subtotal (95% CI)	v				
Total events: 0 (Vitamin D ana Heterogeneity: not applicable	logue), 0 (Dithranol)	Ū			
Total events: 0 (Vitamin D ana Heterogeneity: not applicable Test for overall effect: not appl	logue), 0 (Dithranol) icable			100.0 %	
Total events: 0 (Vitamin D ana	logue), 0 (Dithranol) icable alogue), 45 (Dithranol) hi ² = 8.14, df = 6 (P = 0.23	617	•	100.0 %	-0.03 [-0.06, 0.00]

Analysis 10.8. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 8 Withdrawals due to treatment failure

l Calcipotriol vs. dithranol Berth Jones 1992b	n/N	n/N	H,Random,95%		
1			CI		H,Random,9 Cl
Berth Jones 1992b					
	3/239	3/239	•	77.5 %	0.0 [-0.02, 0.02]
Monastirli 2000	0/35	0/35	+	10.6 %	0.0 [-0.05, 0.05]
Van de Kerkhof 2006	7/54	4/52		2.3 %	0.05 [-0.06, 0.17]
Van der Vleuten 1995	0/10	0/10	-	1.0 %	0.0 [-0.17, 0.17]
Subtotal (95% CI)	338	336	•	91.4 %	0.00 [-0.02, 0.02]
Total events: 10 (Vitamin D anal	logue), 7 (Dithranol)				
Heterogeneity: Tau ² = 0.0; Chi ²	= 1.33, df = 3 (P = 0.72)	$ ^2 = 0.0\%$			
Test for overall effect: $Z = 0.14$	(P = 0.89)				
2 Calcitriol vs. dithranol					
Hutchinson 2000	1/60	2/54	+	8.6 %	-0.02 [-0.08, 0.04]
Subtotal (95% CI)	60	54	•	8.6 %	-0.02 [-0.08, 0.04]
Total events: I (Vitamin D analc	ogue), 2 (Dithranol)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.67$	(P = 0.50)				
3 Tacalcitol vs. dithranol					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D analc	ogue), 0 (Dithranol)				
Heterogeneity: not applicable					
Test for overall effect: not applic	able				
Total (95% CI)	398	390		100.0 %	0.00 [-0.02, 0.02]
Total events: 11 (Vitamin D anal	logue), 9 (Dithranol)				
Heterogeneity: Tau ² = 0.0; Chi ²	= 1.52, df = 4 (P = 0.82)	$ ^2 = 0.0\%$			
Test for overall effect: $Z = 0.06$	(P = 0.95)				
Test for subgroup differences: C	$hi^2 = 0.46, df = 1 (P = 0.5)$	50), l ² =0.0%			

-1 -0.5 0 0.5 I

Analysis 10.9. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 9 Adverse events (local)

Study or subgroup V	Vitamin D analogue	Dithranol	Risk Difference M-	Weight	Risk Difference M-	
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl	
I Calcipotriol vs. dithranol						
Berth Jones 1992b	84/239	127/239	+	12.7 %	-0.18 [-0.27, -0.09]	
Christensen 1999	11/89	23/82	-#-	11.9 %	-0.16 [-0.28, -0.04]	
Grattan 1997 (H)	1/22	11/22		8.9 %	-0.45 [-0.68, -0.23]	
Lister 1997	11/89	23/82		11.9 %	-0.16 [-0.28, -0.04]	
Monastirli 2000	5/35	16/35		9.6 %	-0.31 [-0.52, -0.11]	
Van de Kerkhof 2006	21/53	37/52		10.2 %	-0.32 [-0.50, -0.14]	
Wall 1998	28/161	71/145	-	12.4 %	-0.32 [-0.42, -0.22]	
Subtotal (95% CI)	688	657	•	77.7 %	-0.25 [-0.32, -0.17]	
Hutchinson 2000	3/60	39/54		116%	-0.67 [-0.80 -0.54	
2 Calcitriol vs. dithranol						
Hutchinson 2000	3/60	39/54 54	+ ◆	11.6 %		
Subtotal (95% CI)	60	39/54 54	+ ◆	11.6 %		
Subtotal (95% CI) Total events: 3 (Vitamin D a	60 analogue), 39 (Dithranol)		+ ◆		2	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1	60 analogue), 39 (Dithranol) ble		↓		2	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol	60 analogue), 39 (Dithranol) ole 10.01 (P < 0.00001)	54	+ •	11.6 %	-0.67 [-0.80, -0.54]	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol Farkas 1999	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42	54 17/42	+ •	11.6 %	- 0.67 [-0.80, -0.54]	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol Farkas 1999 Subtotal (95% CI)	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42 42	54	+ + +	11.6 %	- 0.67 [-0.80, -0.54] -0.36 [-0.52, -0.20	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol Farkas 1999 Subtotal (95% CI) Total events: 2 (Vitamin D a	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42 42 analogue), 17 (Dithranol)	54 17/42	+ • •	11.6 %	-0.67 [-0.80, -0.54] -0.67 [-0.80, -0.54] -0.36 [-0.52, -0.20] -0.36 [-0.52, -0.20]	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol Farkas 1999 Subtotal (95% CI) Total events: 2 (Vitamin D a Heterogeneity: not applicab	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42 42 analogue), 17 (Dithranol) ble	54 17/42	+ • •	11.6 %	- 0.67 [-0.80, -0.54]	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol Farkas 1999 Subtotal (95% CI) Total events: 2 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 4	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42 42 analogue), 17 (Dithranol) ble 4.33 (P = 0.000015)	54 17/42 42	+ + +	11.6 % 10.7 % 10.7 %	-0.67 [-0.80, -0.54] -0.36 [-0.52, -0.20] -0.36 [-0.52, -0.20]	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol Farkas 1999 Subtotal (95% CI) Total events: 2 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 4 Total (95% CI)	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42 42 analogue), 17 (Dithranol) ble 4.33 (P = 0.000015) 790	54 17/42	+ + + +	11.6 %	- 0.67 [-0.80, -0.54]	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol Farkas 1999 Subtotal (95% CI) Total events: 2 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 4 Total (95% CI) Total events: 166 (Vitamin D	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42 42 analogue), 17 (Dithranol) ble 4.33 (P = 0.000015) 790 D analogue), 364 (Dithranol)	54 17/42 42 753	+ * -+ *	11.6 % 10.7 % 10.7 %	-0.67 [-0.80, -0.54] -0.36 [-0.52, -0.20] -0.36 [-0.52, -0.20]	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 1 3 Tacalcitol vs. dithranol Farkas 1999 Subtotal (95% CI) Total events: 2 (Vitamin D a Heterogeneity: not applicab Test for overall effect: Z = 4 Total (95% CI) Total events: 166 (Vitamin E Heterogeneity: Tau ² = 0.02;	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42 42 analogue), 17 (Dithranol) ble 4.33 (P = 0.000015) 790 D analogue), 364 (Dithranol) 2; Chi ² = 51.76, df = 8 (P<0.00	54 17/42 42 753	+ + + •	11.6 % 10.7 % 10.7 %	-0.67 [-0.80, -0.54] -0.36 [-0.52, -0.20] -0.36 [-0.52, -0.20]	
Subtotal (95% CI) Total events: 3 (Vitamin D a Heterogeneity: not applicab Test for overall effect: $Z = 1$ 3 Tacalcitol vs. dithranol Farkas 1999 Subtotal (95% CI) Total events: 2 (Vitamin D a Heterogeneity: not applicab Test for overall effect: $Z = 4$ Total (95% CI) Total events: 166 (Vitamin D Heterogeneity: Tau ² = 0.02; Test for overall effect: $Z = 5$	60 analogue), 39 (Dithranol) ble 10.01 (P < 0.00001) 2/42 42 analogue), 17 (Dithranol) ble 4.33 (P = 0.000015) 790 D analogue), 364 (Dithranol) 2; Chi ² = 51.76, df = 8 (P<0.00	54 17/42 42 753 0001); l ² =85%	+ + + •	11.6 % 10.7 % 10.7 %	-0.67 [-0.80, -0.54] -0.36 [-0.52, -0.20] -0.36 [-0.52, -0.20]	

Analysis 10.10. Comparison 10 Vitamin D alone or in combination versus dithranol, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 10 Vitamin D alone or in combination versus dithranol

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D analogue	Dithranol	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,959 Cl
l Calcipotriol vs. dithranol					
Berth Jones 1992b	0/239	1/239	•	81.5 %	0.00 [-0.02, 0.01]
Van der Vleuten 1995	0/35	0/35	+	3.7 %	0.0 [-0.05, 0.05]
Subtotal (95% CI)	274	274	•	85.2 %	0.00 [-0.02, 0.01]
Total events: 0 (Vitamin D a	nalogue), I (Dithranol)				
Heterogeneity: $Tau^2 = 0.0$; (Chi ² = 0.03, df = 1 (P = 0.87);	l ² =0.0%			
Test for overall effect: $Z = 0$	0.70 (P = 0.49)				
2 Calcitriol vs. dithranol					
Hutchinson 2000	0/60	0/54	ŧ	9.5 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	60	54	•	9.5 %	0.0 [-0.03, 0.03]
Total events: 0 (Vitamin D a	nalogue), 0 (Dithranol)				
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
3 Tacalcitol vs. dithranol					
Farkas 1999	0/42	0/42	+	5.3 %	0.0 [-0.05, 0.05]
Subtotal (95% CI)	42	42	+	5.3 %	0.0 [-0.05, 0.05]
Total events: 0 (Vitamin D a	nalogue), 0 (Dithranol)				
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
Total (95% CI)	376	370		100.0 %	0.00 [-0.01, 0.01]
Total events: 0 (Vitamin D a	nalogue), I (Dithranol)				
Heterogeneity: $Tau^2 = 0.0$; (Chi ² = 0.11, df = 3 (P = 0.99);	l ² =0.0%			
Test for overall effect: $Z = 0$	0.64 (P = 0.52)				
Test for subgroup difference	es: $Chi^2 = 0.07$, $df = 2$ (P = 0.9	6), I ² =0.0%			

-I -0.5 0 0.5 I

Analysis 11.1. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 1 IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: I IAGI

Study or subgroup	Vitamin D alone or in combination N	Mean(SD)	Other vitamin D analogue N	Mean(SD)	Std. Mean Difference IV,Random,95% CI	Weight	Std. Mean Difference IV.Random,95% CI
		(ob)		(incari(ob)			
I Calcipotriol vs. calcitriol							
Ji 2008	122	-2.15 (0.76)	124	-2.15 (0.73)		100.0 %	0.0 [-0.25, 0.25]
Subtotal (95% CI)	122		124		+	100.0 %	0.0 [-0.25, 0.25]
Heterogeneity: not applicat	ble						
Test for overall effect: $Z = 0$	0.0 (P = 1.0)						
2 Calcipotriol vs. tacalcitol							
Veien 1997	113	-3.8 (0.9)	113	-3.3 (1.2)		100.0 %	-0.47 [-0.73, -0.21]
Subtotal (95% CI)	113		113		•	100.0 %	-0.47 [-0.73, -0.21]
Heterogeneity: not applicat	ole						
Test for overall effect: $Z = 2$	3.48 (P = 0.00050)						
3 Calcipotriol vs. maxacalci	tol						
Barker 1999 (H)	26	-3.19 (1.27)	26	-3.73 (1.22)		100.0 %	0.43 [-0.12, 0.98]
Subtotal (95% CI)	26		26		-	100.0 %	0.43 [-0.12, 0.98]
Heterogeneity: not applicat	ble						
Test for overall effect: Z =	I.52 (P = 0.13)						
Test for subgroup difference	es: Chi ² = 11.22, d	f = 2 (P = 0.00)	, I ² =82%				
						1	
				-2	-1 0 1	2	

Favours calcipotriol Favours other vitamin D analogue

Analysis 11.2. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 2 TSS

Study or subgroup	Vitamin D alone or in combination		Other vitamin D analogue		Std. Mean Difference	Weight	
	N	Mean(SD)	N	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
l Calcipotriol vs. calcitriol							
Ji 2008	125	1.87 (2.07)	125	2.54 (2.08)	-	41.1 %	-0.32 [-0.57, -0.07]
Subtotal (95% CI) Heterogeneity: not applical	125		125		•	41.1 %	-0.32 [-0.57, -0.07]
Test for overall effect: $Z =$							
2 Calcipotriol vs. tacalcitol							
Veien 1997	145	-5.05 (2.26)	142	-4.03 (2.26)	-	43.3 %	-0.45 [-0.68, -0.22]
Subtotal (95% CI)	145		142		•	43.3 %	-0.45 [-0.68, -0.22]
Heterogeneity: not applical	ble						
Test for overall effect: $Z =$	3.76 (P = 0.00017)						
3 Calcipotriol vs. maxacalci	itol						
Barker 1999 (H)	26	2.8 (2.3)	26	2.4 (3.5)	-	15.6 %	0.13 [-0.41, 0.68]
Subtotal (95% CI)	26		26		+	15.6 %	0.13 [-0.41, 0.68]
Heterogeneity: not applical	ble						
Test for overall effect: Z =	0.48 (P = 0.63)						
Total (95% CI)	296		293		•	100.0 %	-0.31 [-0.55, -0.06]
Heterogeneity: $Tau^2 = 0.02$	2; Chi ² = 3.77, df =	2 (P = 0.15); I ²	=47%				
Test for overall effect: $Z =$	2.47 (P = 0.014)						
Test for subgroup difference	tes: Chi ² = 3.77, df	= 2 (P = 0.15), I	2 =47%				
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				-4	-2 0 2	4	
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Favours calcipotriol Favours other vitamin D analogue

Analysis 11.3. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 3 PASI

Study or subgroup	Vitamin D alone or in combination		Other vitamin D analogue			Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,I	Random,95% Cl	IV,Random,95% CI
l Calcipotriol vs. calcitri	iol						
Bourke 1997	7	4.7 (2.4)	8	8.8 (4.2)			-1.11 [-2.22, 0.01]
2 Calcipotriol vs. tacalci	tol						
3 Calcipotriol vs. maxad	calcitol						
					-10 -5	0 5	10
				F	avours calcipotri	ol Favours c	other vitamin D analogue

Analysis 11.4. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 4 PAGI

Study or subgroup	Vitamin D alone or in combination N	Mean(SD)	Other vitamin D analogue N	Mean(SD)	Std. Mean Difference IV.Random,95% Cl	Std. Mean Difference IV,Random,95% Cl
l Calcipotriol vs. calcitr Ji 2008	riol 125	-2.1 (0.76)	125	-2.13 (0.72)		0.04 [-0.21, 0.29]
2 Calcipotriol vs. tacalc 3 Calcipotriol vs. maxa						_
				F		2 er vitamin D analogue

Analysis 11.5. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D alone or in combination		Other vitamin D analogue	M (CD)	Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	N	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
l Calcipotriol vs. calcitriol							
Bourke 1997	7	4.7 (2.4)	8	8.8 (4.2)		11.0 %	-1.11 [-2.22, 0.01]
Ji 2008	122	-2.15 (0.76)	124	-2.15 (0.73)	+	33.0 %	0.0 [-0.25, 0.25]
Subtotal (95% CI)	129		132		-	44.0 %	-0.41 [-1.46, 0.64]
Heterogeneity: $Tau^2 = 0.44$; Chi ² = 3.61, df =	I (P = 0.06); I ²	=72%				
Test for overall effect: $Z = 0$	0.77 (P = 0.44)						
2 Calcipotriol vs. tacalcitol							
Veien 1997	113	-3.8 (0.9)	113	-3.3 (1.2)	-	32.6 %	-0.47 [-0.73, -0.21]
Subtotal (95% CI)	113		113		•	32.6 %	-0.47 [-0.73, -0.21]
Heterogeneity: not applicab	ble						
Test for overall effect: $Z = 3$	3.48 (P = 0.00050)						
3 Calcipotriol vs. maxacalcit	tol						
Barker 1999 (H)	26	-3.19 (1.27)	26	-3.73 (1.22)	-	23.4 %	0.43 [-0.12, 0.98]
Subtotal (95% CI)	26		26		•	23.4 %	0.43 [-0.12, 0.98]
Heterogeneity: not applicab	ble						
Test for overall effect: $Z = 1$	I.52 (P = 0.13)						
Total (95% CI)	268		271		+	100.0 %	-0.17 [-0.62, 0.27]
Heterogeneity: $Tau^2 = 0.14$; Chi ² = 13.94, df :	= 3 (P = 0.003)	; I ² =78%				
Test for overall effect: $Z = 0$	0.78 (P = 0.44)						
Test for subgroup difference	es: $Chi^2 = 8.33$, df	= 2 (P = 0.02),	$ ^2 = 76\%$				
				1		1	
				_4	4 -2 0 2	4	

Favours calcipotriol Favours other vitamin D analogue

Analysis 11.6. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D alone or in combination	Other vitamin D analogue	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
l Calcipotriol vs. calcitriol					
Bourke 1997	4/12	4/12		2.6 %	0.0 [-0.38, 0.38]
Ji 2008	11/125	8/125	•	85.0 %	0.02 [-0.04, 0.09]
Subtotal (95% CI)	137	137	+	87.6 %	0.02 [-0.04, 0.09]
Total events: 15 (Vitamin D alo	one or in combination), 12	(Other vitamin D analog	gue)		
Heterogeneity: $Tau^2 = 0.0$; Ch	$i^2 = 0.02$, $df = 1$ (P = 0.90)); l ² =0.0%			
Test for overall effect: $Z = 0.7$	I (P = 0.48)				
2 Calcipotriol vs. tacalcitol					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D alor	ne or in combination), 0 (C)ther vitamin D analogue	e)		
Heterogeneity: not applicable					
Test for overall effect: not appl	licable				
3 Calcipotriol vs. maxacalcitol					
Barker 1999 (H)	4/30	4/30		12.4 %	0.0 [-0.17, 0.17]
Subtotal (95% CI)	30	30	+	12.4 %	0.0 [-0.17, 0.17]
Total events: 4 (Vitamin D alor	ne or in combination), 4 (C	ther vitamin D analogue	e)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
Total (95% CI)	167	167	•	100.0 %	0.02 [-0.04, 0.08]
Total events: 19 (Vitamin D ald	one or in combination), 16	(Other vitamin D analog	gue)		
Heterogeneity: $Tau^2 = 0.0$; Ch	$i^2 = 0.08$, df = 2 (P = 0.96)); I ² =0.0%			
Test for overall effect: $Z = 0.66$	6 (P = 0.51)				
Test for subgroup differences:	$Chi^2 = 0.06, df = 1 (P = 0.06)$	80), l ² =0.0%			
			0.5 0 0.5		
				vitamin D analogue	

Analysis 11.7. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D alone or in combination	Other vitamin D analogue	Risk Difference M-	Weight	Risk Difference M
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
l Calcipotriol vs. calcitriol					
Bourke 1997	0/12	0/12	-	5.5 %	0.0 [-0.15, 0.15]
Ji 2008	6/125	2/125	•	63.8 %	0.03 [-0.01, 0.08]
Subtotal (95% CI)	137	137	•	69.4 %	0.03 [-0.01, 0.07]
Total events: 6 (Vitamin D alor	ne or in combination), 2 (C	ther vitamin D analogue	2)		
Heterogeneity: $Tau^2 = 0.0$; Ch	$i^2 = 0.17, df = 1 (P = 0.68)$; l ² =0.0%			
Test for overall effect: $Z = 1.38$	8 (P = 0.17)				
2 Calcipotriol vs. tacalcitol					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D alor	ne or in combination), 0 (C	ther vitamin D analogue	2)		
Heterogeneity: not applicable					
Test for overall effect: not appl	licable				
3 Calcipotriol vs. maxacalcitol					
Barker 1999 (H)	0/30	0/30	+	30.6 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	30	30	•	30.6 %	0.0 [-0.06, 0.06]
Total events: 0 (Vitamin D alor	ne or in combination), 0 (C	ther vitamin D analogue	2)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
Total (95% CI)	167	167	•	100.0 %	0.02 [-0.01, 0.06]
Total events: 6 (Vitamin D alor	ne or in combination), 2 (C	ther vitamin D analogue	2)		
Heterogeneity: $Tau^2 = 0.0$; Ch	$i^2 = 0.79$, df = 2 (P = 0.67)	; l ² =0.0%			
Test for overall effect: $Z = 1.15$	5 (P = 0.25)				
Test for subgroup differences:	$Chi^2 = 0.59, df = 1 (P = 0.59)$	44), I ² =0.0%			
			I -0.5 0 0.5 I		
				vitamin D analogue	

Analysis 11.8. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Vitamin D alone or in combination	Other vitamin D analogue	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
I Calcipotriol vs. calcitriol					
Bourke 1997	1/12	2/12		0.3 %	-0.08 [-0.35, 0.18]
Ji 2008	0/125	0/125	•	93.9 %	0.0 [-0.02, 0.02]
Subtotal (95% CI)	137	137	+	94.2 %	-0.01 [-0.08, 0.07]
Total events: (Vitamin D alc Heterogeneity: Tau ² = 0.00; (, ,	0	e)		
Test for overall effect: $Z = 0.1$	6 (P = 0.87)				
2 Calcipotriol vs. tacalcitol					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D alc	one or in combination), 0 (0	Other vitamin D analogue	e) (e		
Heterogeneity: not applicable					
Test for overall effect: not app	olicable				
3 Calcipotriol vs. maxacalcitol	I				
Barker 1999 (H)	0/30	0/30	+	5.8 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	30	30	•	5.8 %	0.0 [-0.06, 0.06]
Total events: 0 (Vitamin D alc	one or in combination), 0 (0	Other vitamin D analogue	2)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$) (P = 1.0)				
Total (95% CI)	167	167	•	100.0 %	0.00 [-0.02, 0.01]
Total events: I (Vitamin D alc	one or in combination), 2 (C	Other vitamin D analogue	2)		
Heterogeneity: $Tau^2 = 0.0$; Cl	$hi^2 = 0.94, df = 2 (P = 0.62)$	l); l ² =0.0%			
	(P - 0.97)				
Test for overall effect: $Z = 0.0$	(1 - 0.77)				
Test for overall effect: Z = 0.0 Test for subgroup differences:	· · · ·	.90), I ² =0.0%			

Favours calcipotriol

Favours other vitamin D analogue

Analysis 11.9. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D alone or in combination n/N	Other vitamin D analogue n/N	Risk Difference M- H,Random,95% Cl	Weight	Risk Difference H,Random,959 Cl
I Calcipotriol vs. calcitriol					
Ji 2008	14/125	5/125		52.8 %	0.07 [0.01, 0.14]
Subtotal (95% CI)	125	125	•	52.8 %	0.07 [0.01, 0.14]
Total events: 14 (Vitamin D alone	e or in combination), 5 (Other vitamin D analog	ue)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 2.17$ (I	P = 0.030)				
2 Calcipotriol vs. tacalcitol					
Veien 1997	17/145	18/142	-	47.2 %	-0.01 [-0.09, 0.07]
Subtotal (95% CI)	145	142	+	47.2 %	-0.01 [-0.09, 0.07]
Total events: 17 (Vitamin D alone	e or in combination), 18	(Other vitamin D analog	gue)		
Heterogeneity: not applicable					
Test for overall effect: Z = 0.25 (I	P = 0.81)				
3 Calcipotriol vs. maxacalcitol					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D alone	or in combination), 0 (C	Other vitamin D analogue	e)		
Heterogeneity: not applicable					
Test for overall effect: not applica	ble				
Total (95% CI)	270	267	+	100.0 %	0.03 [-0.05, 0.12]
Total events: 31 (Vitamin D alone	e or in combination), 23	(Other vitamin D analog	gue)		
Heterogeneity: $Tau^2 = 0.00$; Chi ²	= 2.69, df = 1 (P = 0.1	0); I ² =63%			
Test for overall effect: $Z = 0.80$ (I	P = 0.42)				
Test for subgroup differences: Ch	$i^2 = 2.56, df = 1 (P = 0)$.), ² =6 %			

Favours calcipotriol

Favours other vitamin D analogue

Analysis 11.10. Comparison 11 Vitamin D alone or in combination versus other vitamin D analogue, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: II Vitamin D alone or in combination versus other vitamin D analogue

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D alone or in combination	Other vitamin D analogue	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
I Calcipotriol vs. calcitriol					
Ji 2008	0/125	0/125	•	42.1 %	0.0 [-0.02, 0.02]
Subtotal (95% CI)	125	125	•	42.1 %	0.0 [-0.02, 0.02]
Total events: 0 (Vitamin D alor	ne or in combination), 0 (O	ther vitamin D analogue)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
2 Calcipotriol vs. tacalcitol					
Veien 1997	0/142	0/145	•	55.3 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	142	145		55.3 %	0.0 [-0.01, 0.01]
Total events: 0 (Vitamin D alor	ne or in combination), 0 (O	ther vitamin D analogue)			
Heterogeneity: not applicable		0,			
Test for overall effect: $Z = 0.0$	(P = 1.0)				
3 Calcipotriol vs. maxacalcitol					
Barker 1999 (H)	0/30	0/30	-	2.6 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	30	30	+	2.6 %	0.0 [-0.06, 0.06]
Total events: 0 (Vitamin D alor	ne or in combination), 0 (O	ther vitamin D analogue)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
Total (95% CI)	297	300	•	100.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Vitamin D alor	ne or in combination), 0 (O	ther vitamin D analogue)			
Heterogeneity: $Tau^2 = 0.0$; Ch	i ² = 0.0, df = 2 (P = 1.00);	l ² =0.0%			
Test for overall effect: $Z = 0.0$	(P = 1.0)				
	$Chi^2 = 0.0, df = 2 (P = 1.00)$	22 12 0.004			

Favours calcipotriol

Favours other vitamin D analogue

Analysis 12.1. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 1 IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: I IAGI

Study or subgroup	Vitamin D alone or in combination		vitamin D and corticosteroid		Std. Mean Difference	Weight	Sto Mea Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% (
I Calcipotriol twice daily	vs. calcipotriol OM, E	MD ON					
Ortonne 1994	80	-3.6 (1)	74	-4.1 (0.76)	-	100.0 %	0.56 [0.23, 0.88
Subtotal (95% CI)	80		74		•	100.0 %	0.56 [0.23, 0.88
Heterogeneity: not applica	able						
Test for overall effect: Z =	= 3.39 (P = 0.00070)						
2 Calcipotriol OD vs. com	nbined calcipotriol +	BMD OD					
Fleming 2010 (H)	74	2.3 (0.81)	150	1.85 (1.05)	-	43.9 %	0.46 [0.18, 0.74
Kaufmann 2002 (H)	480	2.24 (0.9)	490	1.5 (0.9)	•	56.1 %	0.82 [0.69, 0.95
Subtotal (95% CI)	554		640		•	100.0 %	0.66 [0.31, 1.02
Heterogeneity: Tau ² = 0.0 Test for overall effect: Z =		$ (P = 0.02); ^2 =$	=81%				
3 Calcipotriol twice daily	vs. combined calcipo	triol + BMD OD					
Guenther 2002 (H)	227	-3.3 (1.12)	150	-3.59 (1)	-	100.0 %	0.27 [0.06, 0.48
Subtotal (95% CI)	227		150		•	100.0 %	0.27 [0.06, 0.48
Heterogeneity: not applica	able						
Test for overall effect: Z =	. ,						
4 Calcipotriol twice daily				27/00/0		242.04	
Douglas 2002	365	-3.1 (1.09)	369	-3.7 (0.96)		34.3 %	0.58 [0.44, 0.73
	227	-3.3 (1.12)	234	-3.79 (0.97)	•	32.4 %	0.47 [0.28, 0.65
Guenther 2002 (H)	227						
Guenther 2002 (H) Papp 2003 (H)	308	-2.8 (1.21)	301	-3.8 (0.91)	-	33.3 %	0.93 [0.76, 1.10
Papp 2003 (H)		-2.8 (1.21)	301 904	-3.8 (0.91)	•	33.3 % 100.0 %	-
Papp 2003 (H) Subtotal (95% CI)	308 900	~ /	904	-3.8 (0.91)	•		-
	308 900 05; Chi ² = 15.28, df =	~ /	904	-3.8 (0.91)	•		0.93 [0.76, 1.10 0.66 [0.40, 0.93
Papp 2003 (H) Subtotal (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 5 Calcipotriol twice daily	308 900 05; Chi ² = 15.28, df = 4.89 (P < 0.00001) vs. calcipotriol OM, E	= 2 (P = 0.00048	904	-3.8 (0.91)	•	100.0 %	0.66 [0.40, 0.93
Papp 2003 (H) Subtotal (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z =	308 900 05; Chi ² = 15.28, df = : 4.89 (P < 0.00001)	= 2 (P = 0.00048	904	-3.8 (0.91) -3.04 (1.15)	•		0.66 [0.40, 0.93
Papp 2003 (H) Subtotal (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 5 Calcipotriol twice daily	308 900 05; Chi ² = 15.28, df = 4.89 (P < 0.00001) vs. calcipotriol OM, E	= 2 (P = 0.00048 BMV ON	904 1); I ² =87%		•	100.0 %	0.66 [0.40, 0.93
Papp 2003 (H) Subtotal (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 5 Calcipotriol twice daily of Kragballe 1998b Ruzicka 1998	308 900 05; Chi ² = 15.28, df = 4.89 (P < 0.00001) vs. calcipotriol OM, E 172	= 2 (P = 0.00048 BMV ON -2.98 (1.23)	904); I ² =87%	-3.04 (1.15)	•	100.0 % 53.0 % 47.0 %	0.66 [0.40, 0.93 0.05 [-0.16, 0.26 0.53 [0.22, 0.84
Papp 2003 (H) Subtotal (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 5 Calcipotriol twice daily w Kragballe 1998b Ruzicka 1998 Subtotal (95% CI)	308 900 05; Chi ² = 15.28, df = 4.89 (P < 0.00001) vs. calcipotriol OM, E 172 86 258	= 2 (P = 0.00048 MV ON -2.98 (1.23) -3.3 (1.4)	904); l ² =87% 174 78 252	-3.04 (1.15)	•	100.0 % 53.0 % 47.0 %	0.66 [0.40, 0.93 0.05 [-0.16, 0.26 0.53 [0.22, 0.84
Papp 2003 (H) Subtotal (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 5 Calcipotriol twice daily w Kragballe 1998b Ruzicka 1998 Subtotal (95% CI) Heterogeneity: Tau ² = 0.1	308 900 5; Chi ² = 15.28, df = 4.89 (P < 0.00001) vs. calcipotriol OM, E 172 86 258 0; Chi ² = 6.16, df =	= 2 (P = 0.00048 MV ON -2.98 (1.23) -3.3 (1.4)	904); l ² =87% 174 78 252	-3.04 (1.15)	•	100.0 % 53.0 % 47.0 %	0.66 [0.40, 0.93 0.05 [-0.16, 0.26 0.53 [0.22, 0.84
Papp 2003 (H) Subtotal (95% CI) Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 5 Calcipotriol twice daily of Kragballe 1998b	308 900 05; Chi ² = 15.28, df = 4.89 (P < 0.00001) vs. calcipotriol OM, E 172 86 258 0; Chi ² = 6.16, df = = 1.15 (P = 0.25)	= 2 (P = 0.00048 BMV ON -2.98 (1.23) -3.3 (1.4) I (P = 0.01); I ² =	904 174 78 252 =84%	-3.04 (1.15)	•	100.0 % 53.0 % 47.0 %	-

(Continued ...)

Vi	tamin D alone or in	vita	amin D and		Std. Mean		Sto Mea
Study or subgroup	combination	cor	ticosteroid		Difference	Weight	Differenc
	N	Mean(SD)	N	Mean(SD)	IV,Random,95% CI		IV,Random,95% (
Kragballe 1998b	172	-2.98 (1.23)	172	-3.29 (1.09)		100.0 %	0.27 [0.05, 0.48
Subtotal (95% CI)	172		172		•	100.0 %	0.27 [0.05, 0.48
Heterogeneity: not applicable							
Test for overall effect: $Z = 2.46$	` '						
7 Calcipotriol twice daily vs. ca		, ,		,			
Коо 2006	21	3.16 (1.18)	44	2.11 (1.18)		100.0 %	0.88 [0.34, 1.42
Subtotal (95% CI)	21		44		•	100.0 %	0.88 [0.34, 1.42
Heterogeneity: not applicable							
Test for overall effect: $Z = 3.18$	P = 0.0015						
8 Calcipotriol twice daily vs. ca	•	liflucortolone valera					
Subtotal (95% CI)	0		0				Not estimab
Heterogeneity: not applicable							
Test for overall effect: not appl							
9 Calcipotriol OD vs. calcipotr		onide acetonide ON					NT 11
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable Test for overall effect: not appl	anhla						
lest for overall effect. Not appr	ICADIE						
10 Calcipotriol OD vs. combir		,					
Ortonne 2010	202	2.01 (0.95)	206	1.88 (0.94)		100.0 %	0.14 [-0.06, 0.33
Subtotal (95% CI)	202		206		•	100.0 %	0.14 [-0.06, 0.33
Heterogeneity: not applicable							
Test for overall effect: $Z = 1.39$	9 (P = 0.17)						
I Calcitriol twice daily vs. difl	ucortolone vale	rate OM, calcitriol C	N				
Subtotal (95% CI)	0		0				Not estimab
Heterogeneity: not applicable							
Test for overall effect: not appl	icable						
12 Tacalcitol OD vs. combined	calcipotriol + F						
Langley 2011 (H)	163	2.28 (0.89)	171	1.84 (0.93)	-	100.0 %	0.48 [0.26, 0.70
	1(2	~ /	171	()	•	100.0.0/	2
Subtotal (95% CI)	163		171		•	100.0 %	0.48 [0.26, 0.70
Heterogeneity: not applicable Test for overall effect: Z = 4.34	P = 0.000014)					
Test for subgroup differences:		, ,	=61%				
iest for subgroup differences.	20.75, U	0(1 - 0.01), 1 - 0.01)	0170				

Favours vitamin D alone or in combination Favours vitamin D and corticosteroid

Analysis 12.2. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 2 TSS

Study or subgroup	Vitamin D alone or in combination N		ritamin D and orticosteroid N	Mean(SD)		Std. Mean fference om,95% Cl	Std. Mean Difference IV,Random,95% Cl
l Calcipotriol twice dai	ly vs. calcipotriol OM, E	IMD ON					
2 Calcipotriol OD vs. c	ombined calcipotriol +	BMD OD					
3 Calcipotriol twice dai	ly vs. combined calcipo	triol + BMD OD					
Huang 2009	149	-0.071 (0.08)	152	-0.09 (0.07)			0.25 [0.03, 0.48]
4 Calcipotriol twice dai	ly vs. combined calcipo	triol + BMD twice daily					
5 Calcipotriol twice dai	ly vs. calcipotriol OM, E	, MV ON					
6 Calcipotriol twice dai	ly vs. calcipotriol OM, c	lobetasone butyrate Ol	N				
7 Calcipotriol twice dai	ly vs. calcipotriol twice	daily + clobetasol propi	ionate twice daily	,			
8 Calcipotriol twice dai	ly vs. calcipotriol OM, c	liflucortolone valerate (NC				
9 Calcipotriol OD vs. c	alcipotriol OM, fluocino	onide acetonide ON					
10 Calcipotriol OD vs.	combined calcipotriol -	+ hydrocortisone OD					
II Calcitriol twice daily	vs. diflucortolone vale	rate OM, calcitriol ON					
12 Tacalcitol OD vs. co	mbined calcipotriol + E	BMD OD					
					-1 -0.5 (0 0.5 I	

Favours vitamin D alone or in combination

Favours vitamin D and corticosteroid

Analysis 12.3. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 3 PASI

Study or subgroup	Vitamin D alone or in combination	CC	tamin D and orticosteroid		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	N	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
Calcipotriol twice daily v	s. calcipotriol OM, E	BMD ON					
Ortonne 1994	65	-0.74 (0.22)	59	-0.83 (0.16)		100.0 %	0.46 [0.10, 0.82]
ubtotal (95% CI)	65		59		-	100.0 %	0.46 [0.10, 0.82]
eterogeneity: not applica	ble						
est for overall effect: Z =	,						
Calcipotriol OD vs. com			1 47	21 (20)		45.0.0/	0425014.0713
Fleming 2010 (H)	74	4.3 (2.8)	147	3.1 (2.8)	-	45.9 %	0.43 [0.14, 0.71]
Kaufmann 2002 (H)	480	-0.46 (0.31)	490	-0.71 (0.26)		• 54.1 %	0.87 [0.74, 1.01]
ubtotal (95% CI)	554		637		-	- 100.0 %	0.67 [0.23, 1.11]
eterogeneity: $Tau^2 = 0.09$ est for overall effect: Z =		$ (P = 0.005); ^2 =$	=87%				
Calcipotriol twice daily v	· · · · ·	triol + BMD OD					
Guenther 2002 (H)	227	4.2 (3.3)	150	3 (2.5)		29.4 %	0.40 [0.19, 0.61]
Huang 2009	149	-0.7 (0.21)	152	-0.79 (0.18)		26.0 %	0.46 [0.23, 0.69]
Jorizzo 2007	186	-0.43 (0.3)	193	-0.64 (0.3)		29.5 %	0.70 [0.49, 0.91]
Saraceno 2007	73	4.07 (3.33)	74	2.5 (2.5)		15.2 %	0.53 [0.20, 0.86]
ubtotal (95% CI)	635		569		•	100.0 %	0.52 [0.38, 0.67]
eterogeneity: $Tau^2 = 0.0$ est for overall effect: Z =	7.16 (P < 0.00001)						
Calcipotriol twice daily v Douglas 2002	s. combined calcipo 332	trioi + BMD twice 4.4 (3.9)	dally 342	2.5 (2.5)		34.8 %	0.58 [0.43, 0.74]
Guenther 2002 (H)	227	4.2 (3.3)	234	2.7 (2.5)		31.5 %	0.51 [0.33, 0.70]
Papp 2003 (H)	308	-0.49 (0.32)	301	-0.73 (0.25)		- 33.6 %	0.83 [0.67, 1.00]
ubtotal (95% CI)	867		877		•	100.0 %	0.64 [0.46, 0.83]
eterogeneity: $Tau^2 = 0.02$	2; Chi ² = 7.59, df =	2 (P = 0.02); I ² =	74%				
est for overall effect: Z = Calcipotriol twice daily v	· · · · · · · · · · · · · · · · · · ·						
Kragballe 1998b	172 I 172	4.04 (3.39)	174	3.42 (3.05)		52.6 %	0.19 [-0.02, 0.40]
Ruzicka 1998	87	1.9 (1.59)	82	I (0.82)			0.70 [0.39, 1.01]

Favours vitamin D alone or in combination Favours vitamin D and corticosteroid

(Continued . . .)

Study or subgroup	Vitamin D alone or in combination		amin D and rticosteroid		Std. Mean Difference	Weight	Std. Mean Difference
Study of subgroup	N	Mean(SD)	N	Mean(SD)	IV,Random,95% Cl	vveigni	IV,Random,95% CI
Subtotal (95% CI)	259		256			100.0 %	0.43 [-0.07, 0.93]
Heterogeneity: $Tau^2 = 0.11$;	Chi ² = 7.08, df =	$ (P = 0.01); ^2 = 8$	6%				
Test for overall effect: $Z = 1$.	.70 (P = 0.089)						
6 Calcipotriol twice daily vs.	calcipotriol OM, c	lobetasone butyrat	e ON				
Kragballe 1998b	172	4.04 (3.39)	172	3.5 (2.86)		100.0 %	0.17 [-0.04, 0.38]
Subtotal (95% CI)	172		172		-	100.0 %	0.17 [-0.04, 0.38]
Heterogeneity: not applicabl	e						
Test for overall effect: $Z = 1$.59 (P = 0.11)						
7 Calcipotriol twice daily vs.	calcipotriol twice	daily + clobetasol p	propionate tw	rice daily			
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicabl	e						
Test for overall effect: not ap	plicable						
8 Calcipotriol twice daily vs.	calcipotriol OM, c	liflucortolone valer	ate ON				
Salmhofer 2000	. 58	1.9 (1.4)	58	1.8 (1.2)		100.0 %	0.08 [-0.29, 0.44]
Subtotal (95% CI)	58		58			100.0 %	0.08 [-0.29, 0.44]
Heterogeneity: not applicabl Test for overall effect: Z = 0. 9 Calcipotriol OD vs. calcipc Wozel 2001	41 (P = 0.68)	nide acetonide ON 8.94 (5.08)	N 19	6.3 (4.57)		+ 100.0 %	0.53 [-0.11, 1.18]
		0.74 (5.00)		0.5 (1.57)			
Subtotal (95% CI)	19		19			- 100.0 %	0.53 [-0.11, 1.18]
Heterogeneity: not applicabl Test for overall effect: Z = 1.							
10 Calcipotriol OD vs. comb	pined calcipotriol -	- hydrocortisone C	D				
Ortonne 2010	202	4.38 (3.93)	206	4.05 (3.87)		100.0 %	0.08 [-0.11, 0.28]
Subtotal (95% CI)	202		206		-	100.0 %	0.08 [-0.11, 0.28]
Heterogeneity: not applicabl Test for overall effect: Z = 0.							
I I Calcitriol twice daily vs. d	iflucortolone vale	ate OM, calcitriol (NC				
Lee 2007	73	-0.26 (0.33)	69	-0.34 (0.33)	+	100.0 %	0.24 [-0.09, 0.57]
Subtotal (95% CI) Heterogeneity: not applicabl	73		69		-	100.0 %	0.24 [-0.09, 0.57]
Test for overall effect: $Z = 1$.	43 (P = 0.15)						
12 Tacalcitol OD vs. combin	ed calcipotriol + F						
Langley 2011 (H)	163	5.35 (4.44)	171	3.58 (2.98)		100.0 %	0.47 [0.25, 0.69]
Subtotal (95% CI)	163		171		-	100.0 %	0.47 [0.25, 0.69]
Heterogeneity: not applicabl	e						
Test for overall effect: $Z = 4$.)					
Test for subgroup difference			2 =67%				
), .					

Analysis 12.4. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I2 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 4 PAGI

Study or subgroup	Vitamin D alone or in combination		min D and ticosteroid		Std. Mean Difference	Weight	Std Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% Cl		IV,Random,95% C
I Calcipotriol twice daily vs.	calcipotriol OM, B	MD ON					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	2						
Test for overall effect: not ap	plicable						
2 Calcipotriol OD vs. combi	ned calcipotriol +	BMD OD					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	2						
Test for overall effect: not ap	plicable						
3 Calcipotriol twice daily vs.	combined calcipot	riol + BMD OD					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	2						
Test for overall effect: not ap	plicable						
4 Calcipotriol twice daily vs.	combined calcipot	riol + BMD twice d	aily				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	2						
Test for overall effect: not ap	plicable						
5 Calcipotriol twice daily vs.	calcipotriol OM, B	MV ON					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	2						
Test for overall effect: not ap	plicable						
6 Calcipotriol twice daily vs.	calcipotriol OM, cl	obetasone butyrate	ON				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	e						
Test for overall effect: not ap	plicable						
7 Calcipotriol twice daily vs.	calcipotriol twice o	daily + clobetasol pr	opionate twi	e daily			
Коо 2006	21	3.29 (1.3)	44	2.37 (1.3)		14.2 %	0.70 [0.16, 1.23
Subtotal (95% CI)	21		44			14.2 %	0.70 [0.16, 1.23]
Heterogeneity: not applicable						/-	
Test for overall effect: $Z = 2$.							
8 Calcipotriol twice daily vs.	. ,	iflucortolone valerat	te ON				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable			2				
						1	

(Continued \dots)

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Vitamin D alone or in combination	cor	min D and ticosteroid		Std. Mean Difference	Weight	(Continuec Std. Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
Test for overall effect: not ap							
9 Calcipotriol OD vs. calcipo		nide acetonide ON	0				N. 4. 11
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicabl							
Test for overall effect: not ap	plicable						
10 Calcipotriol OD vs. comb	pined calcipotriol +	hydrocortisone OE	C				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicabl	e						
Test for overall effect: not ap	plicable						
I Calcitriol twice daily vs. d	iflucortolone valer	ate OM calcitriol O	N				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable	e						
Test for overall effect: not ap	plicable						
12 Tacalcitol OD vs. combin	ed calcipotriol + B	MD OD					
Langley 2011 (H)	163	2.25 (0.93)	171	1.82 (0.94)		85.8 %	0.46 [0.24, 0.68]
Subtotal (95% CI)	163		171		-	85.8 %	0.46 [0.24, 0.68]
Heterogeneity: not applicabl	e						
Test for overall effect: $Z = 4$.	.14 (P = 0.000035)						
Total (95% CI)	184		215		•	100.0 %	0.49 [0.29, 0.69]
Heterogeneity: $Tau^2 = 0.0$; C	$Chi^2 = 0.67, df = 1$	$(P = 0.4 I); I^2 = 0.09$	6				
Test for overall effect: $Z = 4$.	.80 (P < 0.00001)						
Test for subgroup difference	s: Chi ² = 0.67, df =	$ (P = 0.4), ^2 = 0$.0%				

Favours vitamin D alone or in combination Favours vitamin D and corticosteroid

Analysis 12.5. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

	or in combination		amin D and rticosteroid		Std. Mean Difference	Weight	Std Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% C
I Calcipotriol twice daily v	s. calcipotriol OM, E	BMD ON					
Ortonne 1994	80	-3.6 (1)	74	-4.1 (0.76)	-	100.0 %	0.56 [0.23, 0.88]
Subtotal (95% CI) Heterogeneity: not applical	80		74		*	100.0 %	0.56 [0.23, 0.88]
Test for overall effect: Z =							
2 Calcipotriol OD vs. coml	bined calcipotriol +	BMD OD					
Fleming 2010 (H)	74	2.3 (0.81)	150	1.85 (1.05)	-	43.9 %	0.46 [0.18, 0.74
Kaufmann 2002 (H)	480	2.24 (0.9)	490	1.5 (0.9)		56.1 %	0.82 [0.69, 0.95
Subtotal (95% CI) Heterogeneity: Tau ² = 0.05	554 5; Chi ² = 5.24, df =	(P = 0.02); ² =8	640		•	100.0 %	0.66 [0.31, 1.02]
Test for overall effect: Z =							
3 Calcipotriol twice daily v	s. combined calcipo	triol + BMD OD					
Guenther 2002 (H)	227	-3.3 (1.12)	150	-3.59 (1)	-	27.0 %	0.27 [0.06, 0.48
Huang 2009	149	-0.071 (0.08)	152	-0.09 (0.07)	-	25.8 %	0.25 [0.03, 0.48
Jorizzo 2007	186	-0.43 (0.3)	193	-0.64 (0.3)	-	26.9 %	0.70 [0.49, 0.91
Saraceno 2007	73	4.07 (3.33)	74	2.5 (2.5)	+	20.3 %	0.53 [0.20, 0.86]
Subtotal (95% CI)	635		569		•	100.0 %	0.43 [0.20, 0.66]
Heterogeneity: Tau ² = 0.04	4; Chi ² = 11.46, df =	= 3 (P = 0.01); I ² =	74%				
Test for overall effect: $Z =$	3.68 (P = 0.00023)						
4 Calcipotriol twice daily v			,				
Douglas 2002	365	-3.1 (1.09)	369	-3.7 (0.96)	•	34.3 %	0.58 [0.44, 0.73
Guenther 2002 (H)	227	-3.3 (1.12)	234	-3.79 (0.97)	-	32.4 %	0.47 [0.28, 0.65]
Papp 2003 (H)	308	-2.8 (1.21)	301	-3.8 (0.91)	-	33.3 %	0.93 [0.76, 1.10]
Subtotal (95% CI)	900		904		•	100.0 %	0.66 [0.40, 0.93]
Heterogeneity: $Tau^2 = 0.05$		= 2 (P = 0.00048);	$ ^2 = 87\%$				
Test for overall effect: $Z =$	· · · · · · · · · · · · · · · · · · ·						
5 Calcipotriol twice daily v							
Kragballe 1998b	172	-2.98 (1.23)	174	-3.04 (1.15)	-	53.0 %	0.05 [-0.16, 0.26
Ruzicka 1998	86	-3.3 (1.4)	78	-4 (1.23)	-	47.0 %	0.53 [0.22, 0.84
				-4	-2 0 2	4	

(Continued ...)

	Vitamin D alone or in	vitam	in D and		Std. Mean		St Mea
Study or subgroup	combination N		osteroid N	Mean(SD)	Difference IV,Random,95% CI	Weight	Difference IV,Random,95% (
Subtotal (95% CI)	258	. ,	252		+	100.0 %	0.27 [-0.19, 0.74
Heterogeneity: $Tau^2 = 0.10$;	; Chi ² = 6.16, df =	I (P = 0.0 I); I ² =84%					•
Test for overall effect: $Z = I$.15 (P = 0.25)						
6 Calcipotriol twice daily vs.			N				
Kragballe 1998b	172	-2.98 (1.23)	172	-3.29 (1.09)		100.0 %	0.27 [0.05, 0.48
Subtotal (95% CI)	172		172		•	100.0 %	0.27 [0.05, 0.48
Heterogeneity: not applicab							
Test for overall effect: $Z = 2$. ,						
7 Calcipotriol twice daily vs.					-		0.00 0.004 1.4
Коо 2006	21	3.16 (1.18)	44	2.11 (1.18)		100.0 %	0.88 [0.34, 1.42
Subtotal (95% CI)	21		44		•	100.0 %	0.88 [0.34, 1.42
Heterogeneity: not applicab							
Test for overall effect: Z = 3 8 Calcipotriol twice daily vs.	· · · · · ·	iflucantalana valanata					
Salmhofer 2000	. calcipotriol OP1, c 58	1.9 (1.4)	58	1.8 (1.2)		100.0 %	0.08 [-0.29, 0.44
				110 (112)	T		-
Subtotal (95% CI)	58		58		Ť	100.0 %	0.08 [-0.29, 0.44
Heterogeneity: not applicab Test for overall effect: Z = 0							
P Calcipotriol OD vs. calcip	. ,	nide acetonide ON					
Wozel 2001	19	8.94 (5.08)	19	6.3 (4.57)		100.0 %	0.53 [-0.11, 1.18
Subtotal (95% CI)	19		19		•	100.0 %	0.53 [-0.11, 1.18
Heterogeneity: not applicab			17			100.0 /0	0.95 [*0.11, 1.10
Test for overall effect: $Z = 1$							
10 Calcipotriol OD vs. com	bined calcipotrial -	- hydrocortisone OD					
Ortonne 2010	202	2.01 (0.95)	206	1.88 (0.94)	+	100.0 %	0.14 [-0.06, 0.33
		2.01 (0.70)			T		-
Subtotal (95% CI) Heterogeneity: not applicab	202		206		ľ	100.0 %	0.14 [-0.06, 0.33
Test for overall effect: $Z = 1$							
	, <i>,</i>						
I I Calcitriol twice daily vs. o Lee 2007	3iflucortolone valer 73	-0.26 (0.33)	1 69	-0.34 (0.33)		100.0 %	0.24 [-0.09, 0.57
		-0.26 (0.55)		-0.55) +0.55)			
Subtotal (95% CI)	73		69		•	100.0 %	0.24 [-0.09, 0.57
Heterogeneity: not applicab Test for overall effect: Z = 1							
	· · · ·						
12 Tacalcitol OD vs. combir						100.0.0/	
Langley 2011 (H)	163	2.28 (0.89)	171	1.84 (0.93)		100.0 %	0.48 [0.26, 0.70
Subtotal (95% CI) Heterogeneity: not applicab	163 le		171		•	100.0 %	0.48 [0.26, 0.70
Test for overall effect: $Z = 4$							
Test for subgroup difference	es: Chi ² = 23.13, df	= (P = 0.02), ² =	52%				

Analysis 12.6. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D alone or in combination	vitamin D and corticosteroid	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
I Talcipotriol vs. calcipotriol ar	nd corticosteroid				
Douglas 2002	37/369	28/372	-	11.3 %	0.03 [-0.02, 0.07]
Fleming 2010 (H)	6/79	13/162		3.7 %	0.00 [-0.08, 0.07]
Guenther 2002 (H)	23/231	30/389	+	8.6 %	0.02 [-0.02, 0.07]
Huang 2009	7/160	6/160	+	10.0 %	0.01 [-0.04, 0.05]
Kaufmann 2002 (H)	39/480	13/490	-	22.2 %	0.05 [0.03, 0.08]
Коо 2006	0/21	0/44	-	4.0 %	0.0 [-0.07, 0.07]
Kragballe 1998b	19/174	23/351	-	6.8 %	0.04 [-0.01, 0.10]
Ortonne 2010	34/202	27/206		4.0 %	0.04 [-0.03, 0.11]
Papp 2003 (H)	27/308	16/304	-	11.5 %	0.04 [-0.01, 0.08]
Ruzicka 1998	5/87	6/82	<u> </u>	3.5 %	-0.02 [-0.09, 0.06]
Salmhofer 2000	5/63	5/63		2.2 %	0.0 [-0.09, 0.09]
Saraceno 2007	14/75	3/75		2.0 %	0.15 [0.05, 0.25]
Wozel 2001	0/19	0/19	—	2.1 %	0.0 [-0.10, 0.10]
Subtotal (95% CI)	2268	2717	•	91.8 %	0.03 [0.01, 0.05]
Total events: 216 (Vitamin D a Heterogeneity: Tau ² = 0.00; C Test for overall effect: Z = 3.64 2 Calcitriol vs. calcitriol and co Lee 2007	$Chi^2 = 13.84, df = 12 (P + 4 (P = 0.00028))$,	osteroid)	2.5 %	0.01 [-0.08, 0.10]
Subtotal (95% CI)	73	69	•	2.5 %	0.01 [-0.08, 0.10]
Fotal events: 6 (Vitamin D alor Heterogeneity: not applicable Fest for overall effect: $Z = 0.2$,	(vitamin D and corticoster	oid)		
		-(0.5 -0.25 0 0.25 0.5		
		Favours vitamin D alone or in	combination Favours vitamin	D and corticosteroid	(Continued

Topical treatments for chronic plaque psoriasis (Review)

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(... Continued)

					(Continued)
Study or subgroup	Vitamin D alone or in combination n/N	vitamin D and corticosteroid n/N	Risk Difference M- H,Random,95%	Weight	Risk Difference H,Random,95%
3 Tacalcitol vs. calcipotriol and		n/IN	C		Cl
Langley 2011 (H)	21/184	2/ 83		5.6 %	0.05 [-0.01, 0.11]
Subtotal (95% CI)	184	183	•	5.6 %	0.05 [-0.01, 0.11]
Total events: 21 (Vitamin D al			oroid)	J.U /0	0.09[-0.01, 0.11]
,	,	2 (vitamin D and corticost	eroid)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 1.6$	· ,	20(0		100.0.0/	
Total (95% CI)	2525	2969		100.0 %	0.03 [0.02, 0.05]
Total events: 243 (Vitamin D			osteroid)		
Heterogeneity: $Tau^2 = 0.00$; (Chi ² = 14.41, df = 14 (P =	= 0.42); I ² =3%			
Test for overall effect: $Z = 4.3$	36 (P = 0.000013)				
Test for subgroup differences:	$Chi^2 = 0.60, df = 2 (P =$	0.74), l ² =0.0%			
0	× ×	,			
		-(0.5 -0.25 0 0.25 0.5		

Favours vitamin D alone or in combination Favours vitamin D and corticosteroid

Analysis 12.7. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D alone or in combination n/N	vitamin D and corticosteroid n/N	Risk Difference M- H,Random,95% Cl	Weight	Risk Difference H,Random, Cl
Guenther 2002 (H)	6/227	1/386	-	14.2 %	0.02 [0.00, 0.05
Huang 2009	1/160	1/160	+	18.3 %	0.0 [-0.02, 0.02
Kaufmann 2002 (H)	15/480	3/490	-=	18.6 %	0.03 [0.01, 0.04
Коо 2006	0/21	0/44		2.0 %	0.0 [-0.07, 0.07
Kragballe 1998b	8/174	6/351		7.3 %	0.03 [-0.01, 0.06
Ortonne 1994	6/97	3/91		2.6 %	0.03 [-0.03, 0.09
Ortonne 2010	19/202	6/206		4.3 %	0.06 [0.02, 0.11
Ruzicka 1998	1/87	1/82	-	7.7 %	0.00 [-0.03, 0.03
Salmhofer 2000	1/63	0/63		4.9 %	0.02 [-0.03, 0.06
Saraceno 2007	2/75	0/75		4.7 %	0.03 [-0.02, 0.07
Wozel 2001	0/19	0/19		1.1 %	0.0 [-0.10, 0.10
Subtotal (95% CI)	1605	1967	•	1.1 % 85.8 %	0.0 [-0.10, 0.10 0.02 [0.01, 0.03
Subtotal (95% CI) Total events: 59 (Vitamin D a Heterogeneity: Tau ² = 0.00; Test for overall effect: Z = 3. 2 Calcitriol vs. calcitriol and c	1605 alone or in combination), 2 Chi ² = 14.91, df = 10 (P = 01 (P = 0.0026) corticosteroid	1967 21 (vitamin D and corticost = 0.14); I ² =33%		85.8 %	0.02 [0.01, 0.03
Subtotal (95% CI) Fotal events: 59 (Vitamin D a Heterogeneity: Tau ² = 0.00; Test for overall effect: Z = 3. 2 Calcitriol vs. calcitriol and o Lee 2007	1605 alone or in combination), 2 Chi ² = 14.91, df = 10 (P = 01 (P = 0.0026) corticosteroid 2/73	1967 21 (vitamin D and corticost = 0.14); I ² =33% 0/69		85.8 % 4.4 %	0.02 [0.01, 0.03
Subtotal (95% CI) Total events: 59 (Vitamin D a Heterogeneity: Tau ² = 0.00; Test for overall effect: Z = 3. 2 Calcitriol vs. calcitriol and c	1605 alone or in combination), 2 Chi ² = 14.91, df = 10 (P = 01 (P = 0.0026) corticosteroid 2/73 73 one or in combination), 0 e .18 (P = 0.24)	1967 21 (vitamin D and corticost = 0.14); I ² =33% 0/69 69	eroid)	85.8 %	0.02 [0.01, 0.03
Subtotal (95% CI) Fotal events: 59 (Vitamin D a Heterogeneity: Tau ² = 0.00; Test for overall effect: Z = 3. 2 Calcitriol vs. calcitriol and c Lee 2007 Subtotal (95% CI) Fotal events: 2 (Vitamin D al Heterogeneity: not applicable Fest for overall effect: Z = 1.	1605 alone or in combination), 2 Chi ² = 14.91, df = 10 (P = 01 (P = 0.0026) corticosteroid 2/73 73 one or in combination), 0 e .18 (P = 0.24)	1967 21 (vitamin D and corticost = 0.14); I ² =33% 0/69 69	eroid)	85.8 % 4.4 %	0.02 [0.01, 0.03
Subtotal (95% CI) otal events: 59 (Vitamin D a deterogeneity: Tau ² = 0.00; est for overall effect: $Z = 3$. Calcitriol vs. calcitriol and o Lee 2007 Subtotal (95% CI) otal events: 2 (Vitamin D al- deterogeneity: not applicable est for overall effect: $Z = 1$. Tacalcitol vs. calcipotriol an	1605 alone or in combination), 2 Chi² = 14.91, df = 10 (P = 01 (P = 0.0026) corticosteroid 2/73 73 one or in combination), 0 e .18 (P = 0.24) ad corticosteroid	1967 21 (vitamin D and corticost = 0.14); I ² =33% 0/69 69 (vitamin D and corticoster	eroid)	85.8 % 4.4 % 4.4 %	0.02 [0.01, 0.03 0.03 [-0.02, 0.07 0.03 [-0.02, 0.07

(Continued . . .)

(... Continued)

Study or subgroup	Vitamin D alone or in combination n/N	vitamin D and corticosteroid n/N	Risk Difference H,Random,9 Cl	Weight 5%	Risk Difference H,Random,95% Cl
Total (95% CI)	1862	2219	•	100.0 %	0.02 [0.01, 0.03]
Total events: 65 (Vitamin D a	lone or in combination), 2	4 (vitamin D and corticost	eroid)		
Heterogeneity: $Tau^2 = 0.00$; (Chi ² = 15.27, df = 12 (P =	= 0.23); I ² =21%			
Test for overall effect: $Z = 3.3$	36 (P = 0.00077)				
Test for subgroup differences	: $Chi^2 = 0.94$, $df = 2$ (P =	0.62), l ² =0.0%			
		-C	.2 -0.1 0 0	0.1 0.2	
		Favours vitamin D alone or in	combination Fav	ours vitamin D and corticosteroid	

Analysis 12.8. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 8 Withdrawals due to treatment failure

or in combination	vitamin D and corticosteroid	Risk Difference M-	Weight	Risk Difference M-
n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
nd corticosteroid				
2/227	1/386	-	53.0 %	0.01 [-0.01, 0.02]
0/21	0/44	-	1.9 %	0.0 [-0.07, 0.07]
3/174	2/351	-	21.1 %	0.01 [-0.01, 0.03]
8/202	4/206	-	8.5 %	0.02 [-0.01, 0.05]
0/63	0/63	+	9.9 %	0.0 [-0.03, 0.03]
2/75	1/75	+	4.6 %	0.01 [-0.03, 0.06]
0/19	0/19		1.0 %	0.0 [-0.10, 0.10]
781	1144	•	100.0 %	0.01 [0.00, 0.02]
one or in combination), 8	(vitamin D and corticoste	roid)		
	n/N nd corticosteroid 2/227 0/21 3/174 8/202 0/63 2/75 0/19 781	n/N n/N nd corticosteroid 2/227 1/386 0/21 0/44 3/174 2/351 8/202 4/206 0/63 0/63 2/75 1/75 0/19 0/19 781 1144 one or in combination), 8 (vitamin D and corticoster	M- H,Random,95% CI n/N n/N CI nd corticosteroid - - 0/21 0/44 - 3/174 2/351 - 8/202 4/206 - 0/63 0/63 - 0/19 0/19 -	M- H,Random,95% Cl M- H,Random,95% Cl ad corticosteroid 53.0 % 2/227 1/386 53.0 % 0/21 0/44 1.9 % 3/174 2/351 21.1 % 8/202 4/206 8.5 % 0/63 0/63 9.9 % 2/75 1/75 4.6 % 0/19 0/19 1.0 % 781 1144 100.0 %

Favours vitamin D alone or in combination

Favours vitamin D and corticosteroid

(Continued ...)

Topical treatments for chronic plaque psoriasis (Review)

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(... Continued)

					(Continued)
Study or subgroup	Vitamin D alone or in combination	vitamin D and corticosteroid	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
Heterogeneity: $Tau^2 = 0.0$; C	$Chi^2 = 1.36, df = 6 (P = 0.9)$	97); I ² =0.0%			
Test for overall effect: $Z = I$.	.65 (P = 0.099)				
2 Calcitriol vs. calcitriol and o	corticosteroid				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D al	one or in combination), 0	vitamin D and corticosteroi	d)		
Heterogeneity: not applicable	e				
Test for overall effect: not ap	plicable				
3 Tacalcitol vs. calcipotriol ar	nd corticosteroid				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D al	one or in combination), 0	vitamin D and corticosteroi	d)		
Heterogeneity: not applicable	e				
Test for overall effect: not ap	plicable				
Total (95% CI)	781	1144	•	100.0 %	0.01 [0.00, 0.02]
Total events: 15 (Vitamin D a	alone or in combination), 8	(vitamin D and corticostere	(bid		
Heterogeneity: $Tau^2 = 0.0$; C	$Chi^2 = 1.36, df = 6 (P = 0.9)$	97); I ² =0.0%			
Test for overall effect: $Z = 1$.	.65 (P = 0.099)				
Test for subgroup differences	s: Not applicable				
		-0.			
		Favours vitamin D alone or in (complination Favours vitami	n D and corticosteroid	

Analysis 12.9. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D alone or in combination	vitamin D and corticosteroid	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random, C
Calcipotriol vs. calcipotriol ar	nd corticosteroid				
Douglas 2002	44/369	30/372	-	15.4 %	0.04 [0.00, 0.08]
Fleming 2010 (H)	8/79	12/160		5.0 %	0.03 [-0.05, 0.10]
Guenther 2002 (H)	45/227	40/386	-#-	8.3 %	0.09 [0.03, 0.15]
Huang 2009	25/160	17/160		5.6 %	0.05 [-0.02, 0.12]
Kaufmann 2002 (H)	54/480	29/490	-	22.1 %	0.05 [0.02, 0.09]
Kragballe 1998b	54/173	71/347		4.7 %	0.11 [0.03, 0.19]
Ortonne 1994	24/94	11/88		2.5 %	0.13 [0.02, 0.24
Ortonne 2010	63/200	40/204		4.3 %	0.12 [0.03, 0.20
Papp 2003 (H)	53/308	30/304	-=-	10.2 %	0.07 [0.02, 0.13
Ruzicka 1998	3/87	6/82		3.5 %	0.08 [-0.02, 0.17
Salmhofer 2000	6/63	8/63		2.6 %	-0.03 [-0.14, 0.08
Saraceno 2007	8/75	7/75		3.4 %	0.01 [-0.08, 0.11
Wozel 2001	1/19	1/19		1.6 %	0.0 [-0.14, 0.14
ubtotal (95% CI)	2334	2750	•	89.1 %	0.06 [0.04, 0.08]
otal events: 398 (Vitamin D al leterogeneity: Tau ² = 0.00; CP est for overall effect: Z = 5.92 Calcitriol vs. calcitriol and cor	$hi^2 = 13.26, df = 12 (P = 12)$		osteroid)		
Lee 2007	8/67	1/64		4.4 %	0.10 [0.02, 0.19
ubtotal (95% CI) otal events: 8 (Vitamin D alon leterogeneity: not applicable est for overall effect: Z = 2.44 Tacalcitol vs. calcipotriol and	(P = 0.015)	64 (vitamin D and corticostered	pid)	4.4 %	0.10 [0.02, 0.19]
Langley 2011 (H)	32/184	16/182		6.5 %	0.09 [0.02, 0.15]
ubtotal (95% CI)	184	182	•	6.5 %	0.09 [0.02, 0.15]

(Continued . . .)

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Study or subgroup	Vitamin D alone or in combination	vitamin D and corticosteroid	Risk Difference Mi	e Weight	Risk Difference M- H,Random,95%
	n/N	n/N	H,Random,' C		H,Random,95% Cl
Total events: 32 (Vitamin D ald	one or in combination), I	6 (vitamin D and corticoste	eroid)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 2.46$	6 (P = 0.014)				
Total (95% CI)	2585	2996	•	100.0 %	0.06 [0.05, 0.08]
Total events: 438 (Vitamin D a	lone or in combination),	319 (vitamin D and cortico	osteroid)		
Heterogeneity: $Tau^2 = 0.00$; C	hi ² = 14.60, df = 14 (P =	= 0.4); ² =4%			
Test for overall effect: $Z = 6.94$	4 (P < 0.00001)				
Test for subgroup differences:	$Chi^2 = 1.46$, df = 2 (P =	0.48), l ² =0.0%			
		-(0.5 -0.25 0	0.25 0.5	
		Favours vitamin D alone or in	combination Fa	vours vitamin D and corticosteroid	

Analysis 12.10. Comparison 12 Vitamin D alone or in combination versus vitamin D + corticosteroid, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 12 Vitamin D alone or in combination versus vitamin D + corticosteroid

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D alone or in combination	vitamin D and corticosteroid	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Calcipotriol vs. calcipotriol a	nd corticosteroid				
Douglas 2002	0/369	0/372	+	56.2 %	0.0 [-0.01, 0.01]
Huang 2009	3/160	3/160		1.8 %	0.0 [-0.03, 0.03]
Papp 2003 (H)	0/308	0/304	+	38.4 %	0.0 [-0.01, 0.01]
Ruzicka 1998	/87	7/82		0.2 %	0.04 [-0.05, 0.13]
Salmhofer 2000	0/63	0/63		1.7 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	987	981		98.2 %	0.00 [0.00, 0.00]
Test for overall effect: Z = 0.0- 2 Calcitriol vs. calcitriol and co	orticosteroid	0/64		18%	
	. ,				
Lee 2007	0/67	0/64		1.8 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	67	64	-	1.8 %	0.0 [-0.03, 0.03]
Total events: 0 (Vitamin D alor	ne or in combination), 0	(vitamin D and corticosterc	pid)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
3 Tacalcitol vs. calcipotriol and					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D alor	ne or in combination), 0	(vitamin D and corticosterc	pid)		
Heterogeneity: not applicable					
Test for overall effect: not app					
Total (95% CI)	1054	1045		100.0 %	0.00 [0.00, 0.00]
Total events: 14 (Vitamin D alo	one or in combination), I	0 (vitamin D and corticoste	eroid)		
	$i^2 = 3.32$, df = 5 (P = 0.6)	65); I ² =0.0%			
Heterogeneity: Tau ² = 0.0; Ch					
Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 0.0	4 (P = 0.97)				
8 ,	(/	1.00), I ² =0.0%			

Favours vitamin D alone or in combination Favours vitamin D and corticosteroid

Analysis 13.1. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 1 IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

Outcome: I IAGI

Study or subgroup	Vitamin D regimen		Complex regimen		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	N	Mean(SD)	IV,Random,95% Cl		IV,Random,95% CI
I Calcipotriol (12 wks) vs	s. combined calcipotr	iol + BMD (8 w	vks); then calcipotri	ol (4 wks)	_		
Kragballe 2004	283	1.62 (0.86)	294	1.73 (0.92)		100.0 %	-0.12 [-0.29, 0.04]
Subtotal (95% CI)	283		294			100.0 %	-0.12 [-0.29, 0.04]
Heterogeneity: not applic	able						
Test for overall effect: Z =	= 1.48 (P = 0.14)						
2 Calcipotriol (12 wks) vs	s. combined calcipotr	iol + BMD (4 w	vks); then calcipotri	ol (8 wks)			
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	able						
Test for overall effect: not	t applicable						
3 Calcipotriol (12 wks) vs	s. combined calcipotr	iol + BMD (4 w	vks); then calcipotri	ol (w/dy) % com	pined calcipotriol + BMD	D (w/e) (8 wks)	
Kragballe 2004	283	1.62 (0.86)	302	1.51 (0.89)	+	100.0 %	0.13 [-0.04, 0.29]
Subtotal (95% CI)	283		302		-	100.0 %	0.13 [-0.04, 0.29]
Heterogeneity: not applic	able						
Test for overall effect: Z =	= 1.51 (P = 0.13)						
4 Calcipotriol (6 wks) vs.	clobetasol propionat	e (2 wks); then	calcipotriol (4 wks))			
Austad 1998	46	-3.17 (0.82)	46	-3.67 (0.84)		→ 100.0 %	0.60 [0.18, 1.02]
Subtotal (95% CI)	46		46			100.0 %	0.60 [0.18, 1.02]
Heterogeneity: not applic	able						
Test for overall effect: Z =	= 2.80 (P = 0.0051)						
5 Calcipotriol (6 wks) vs.	calcipotriol OM, fluc	cinonide acetor	nide ON (2 wks); tl	hen calcipotriol t	wice daily (4 wks)		
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	able						
Test for overall effect: not	t applicable						
6 Calcipotriol (6 wks) vs.	halometasone OM,	calcipotriol ON	(2 wks); then calci	potriol twice dail	y (w/dy), halometasone	(w/e) (2 wks); tł	nen calcipotriol twice daily
(2wks)							
Yang 2009	36	-1.86 (0.93)	40	-2.23 (0.86)		• 100.0 %	0.41 [-0.05, 0.86]
Subtotal (95% CI)	36		40			100.0 %	0.41 [-0.05, 0.86]
Heterogeneity: not applic	able						
Heterogeneity: not applic Test for overall effect: Z =							
Test for overall effect: Z =	= 1.76 (P = 0.078)	(2 to 4 wks); t	hen calcipotriol twi	ice daily (to wk 1	2) vs. calcitriol ON, clob	etasol propiona	te OM (2 to 4 wks); then
Test for overall effect: Z = 7 Calcipotriol ON, clober calcitriol twice daily (to	= 1.76 (P = 0.078) tasol propionate OM wk 12)	. ,	•	, , ,	2) vs. calcitriol ON, clob		· · ·
Test for overall effect: Z = 7 Calcipotriol ON, clobe	= 1.76 (P = 0.078) tasol propionate OM	(2 to 4 wks); t -4.69 (1.3)	•	ice daily (to wk ∣ -4.44 (1.28) ←	2) vs. calcitriol ON, clob	etasol propiona 100.0 %	· · ·
Test for overall effect: Z = 7 Calcipotriol ON, clober calcitriol twice daily (to	= 1.76 (P = 0.078) tasol propionate OM wk 12)	. ,	•	-4.44 (1.28) ←		100.0 %	te OM (2 to 4 wks); then -0.19 [-0.54, 0.16]
Test for overall effect: Z = 7 Calcipotriol ON, clober calcitriol twice daily (to	= 1.76 (P = 0.078) tasol propionate OM wk 12)	. ,	•	, , ,	-0.25 0 0.25		. ,

Study or subgroup	Vitamin D regimen		plex regimen		Std. Mean Difference	Weight	(Continue Std Mear Difference
	N	Mean(SD)	N	Mean(SD)	IV,Random,95% Cl		IV,Random,95% C
Subtotal (95% CI)			61			100.0 %	-0.19 [-0.54, 0.16
Heterogeneity: not appli							
Test for overall effect: Z	· /						
	+ BMD (4 wks); then	placebo ointment t	wice daily (8 v	vks) vs. combine	d calcipotriol + BMD (4 w	/ks); then calcip	otriol ointment twice dail
(8 wks) White 2006 (P)	376	2.49 (0.99)	383	2.23 (0.96)		100.0 %	0.27 [0.12, 0.41
Subtotal (95% CI)	376		383		-	100.0 %	0.27 [0.12, 0.41
Heterogeneity: not appli Test for overall effect: Z							
	· · · · ·	placebo ointment t	wice daily (8)	uks) vs. combine	d calcipotriol + BMD (4 v	uks): then calcir	otrial (w/dv)+ combined
calcipotriol + BMD (w. White 2006 (P)	. ,	2.49 (0.99)	377	1.98 (0.99)		• 100.0 %	0.51 [0.37, 0.66
		2.17 (0.77)					-
Subtotal (95% CI)			377			100.0 %	0.51 [0.37, 0.66
Heterogeneity: not appli							
Test for overall effect: Z	= 6.95 (P < 0.00001)						
10 Combined calcipotric	ol + BMD (4 wks); the	en calcipotriol ointm	nent twice dail	y (8 wks) vs. coi	mbined calcipotriol + BM	D (4 wks); the	n calcipotriol (w/dy)+
combined calcipotriol	+ BMD (w/e) (8 wks)					
White 2006 (H)	383	2.23 (0.96)	377	1.98 (0.99)		100.0 %	0.26 [0.11, 0.40
Subtotal (95% CI)	383		377		-	100.0 %	0.26 [0.11, 0.40
Heterogeneity: not appli	able						•
Test for overall effect: Z	= 3.52 (P = 0.00044)						
			N				
	I + BMD (8 wks); the	n calcipotriol (4 wks	s) vs. combine	d calcipotriol +	BMD (4 wks); then calcipo	otriol (w/dy) %	combined calcipotriol +
BMD (w/e) (8 wks) Kragballe 2004	294	1.73 (0.92)	302	1.51 (0.89)		100.0 %	0.24 [0.08, 0.40
0							-
Subtotal (95% CI)			302			100.0 %	0.24 [0.08, 0.40
Heterogeneity: not appli							
Test for overall effect: Z	= 2.95 (P = 0.0032)						
12 Tacalcitol (8 wks) vs.	combined calcipotriol	+ BMD (4 wks); the	n calcipotriol	(4 wks)			
Ortonne 2004	247	-2.45 (1.31)	246	-3.16 (1.33)		100.0 %	0.54 [0.36, 0.72
Subtotal (95% CI)	247		246			100.0 %	0.54 [0.36, 0.72
Heterogeneity: not appli			240			100.0 /0	0.71 [0.30, 0.72
Test for overall effect: Z							
Test for subgroup differe	· · · · ·	$= 9 (P = 0.00) l^2 =$	83%				
iest for subgroup differe		∑ (i = 0.00), i =	00/0				
				-0.5	-0.25 0 0.25	0.5	

Analysis 13.2. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

Outcome: 2 TSS

Study or subgroup	Vitamin D regimen	C	Complex regimen		Std. Mean Difference	Weight	Std. Mean Difference
study of subgroup	N	Mean(SD)	ompiex regimen N	Mean(SD)	IV.Random.95% CI	vveigin	IV.Random.95% CI
Calcipotriol (12 wks) vs	s combined calcipotric	()	s): then calcipotriol	()			,
Subtotal (95% CI)	0		0	(1 110)			Not estimable
Heterogeneity: not applic	-		Ū				1.00 0000000000000000000000000000000000
Test for overall effect: not							
2 Calcipotriol (12 wks) vs		ol + BMD (4 wks	s): then calcipotriol	(8 wks)			
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	able						
Test for overall effect: not							
3 Calcipotriol (12 wks) vs		ol + BMD (4 wks	s); then calcipotriol	(w/dy) % comb	pined calcipotriol + BMD	(w/e) (8 wks)	
Subtotal (95% CI)	0	,	0	())		. , . ,	Not estimable
Heterogeneity: not applic	able						
Test for overall effect: not	applicable						
4 Calcipotriol (6 wks) vs.		(2wks); then ca	lcipotriol (4 wks)				
Austad 1998	46	2.5 (1.3)	46	1.7 (1.2)		100.0 %	0.63 [0.21, 1.05]
Subtotal (95% CI)	46		46		•	100.0 % 0	.63 [0.21, 1.05]
Heterogeneity: not applic	able						
Test for overall effect: Z =							
5 Calcipotriol (6 wks) vs.	· · · · · ·	inonide acetonic	le ON (2 wks): the	n calcipotriol tv	vice daily (4 wks)		
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	able						
Test for overall effect: not							
6 Calcipotriol (6 wks) vs.		lcipotriol ON (2	wks); then calcipot	riol twice daily	(w/dy), halometasone (w	//e) (2 wks); then ca	lcipotriol twice daily (2
wks)							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	able						
Test for overall effect: not	applicable						
7 Calcipotriol ON, clobe	tasol propionate OM	(2 to 4 wks); the	n calcipotriol twice	daily (to wk 12	2) vs. calcitriol ON, clobe	etasol propionate O	M (2 to 4 wks); then
calcitriol twice daily (to	wk 12)						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applic	able						
Test for overall effect: not	applicable						
8 Combined calcipotriol	+ BMD (4 wks); then	placebo ointmer	nt twice daily (8 wk	s) vs. combined	calcipotriol + BMD (4 v	vks); then calcipotric	ol ointment twice daily
(8 wks) Subtotal (95% CI)	0		0				Not estimable
(99% CI)	U		0			1	Not estimable
				-2	-I 0 I	2	
				Favours vitamin	D regimen Favours of	complex regimen	1-
							(Continued

Study or subgroup	Vitamin D regimen N	Compl Mean(SD)	ex regimen N Mea	an(SD)		Std. Mean Irence n,95% Cl	Weight	(Continued Std. Mean Difference IV.Random,95% CI
Heterogeneity: not appli	cable	. ,		()				
Test for overall effect: no	ot applicable							
9 Combined calcipotriol calcipotriol + BMD (w	()	olacebo ointment twic	e daily (8 wks) vs.	combined ca	lcipotriol +	BMD (4 wks)); then calcipotrio	l (w/dy) + combined
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applie Test for overall effect: no								
10 Combined calcipotric combined calcipotrio	ol + BMD (4 wks); the l + BMD (w/e) (8 wks)	n calcipotriol ointmen	t twice daily (8 wk	s) vs. combii	ned calcipot	riol + BMD ((4 wks); then calc	ipotriol (w/dy) +
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not appli	cable							
Test for overall effect: no	ot applicable							
II Combined calcipotric BMD (w/e) (8 wks)	bl + BMD (8 wks); ther	a calcipotriol (4 wks) v	vs. combined calcip	otriol + BMI	D (4 wks); t	hen calcipotri	iol (w/dy) % com	bined calcipotriol +
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not appli	cable							
Test for overall effect: no	ot applicable							
12 Tacalcitol (8 wks) vs.	combined calcipotriol +	- BMD (4 wks); then a	alcipotriol (4 wks)					
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not appli	cable							
Test for overall effect: no	ot applicable							
Test for subgroup differe	nces: Not applicable							
				1			í.	
				-2	- I 0	1 2	2	
			Favo	urs vitamin D	regimen	Favours com	plex regimen	

Analysis 13.3. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

Outcome: 3 PASI

Study or subgroup Vitamin	D regimen N	Con Mean(SD)	nplex regimen N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
I Calcipotriol (12 wks) vs. combin	ed calcipotric	l + BMD (8 wks); ti	hen calcipotriol	(4 wks)			
Kragballe 2004	327 -	0.659 (0.331)	322 -	-0.65 (0.317)		100.0 %	-0.04 [-0.19, 0.11]
Subtotal (95% CI)	327		322		•	100.0 %	-0.04 [-0.19, 0.11]
Heterogeneity: not applicable							
Test for overall effect: $Z = 0.51$ (P	= 0.61)						
2 Calcipotriol (12 wks) vs. combin	ed calcipotric	l + BMD (4 wks); tl	hen calcipotriol	(8 wks)			
Saraceno 2007	71	3.04 (3.76)	72	2.11 (2.56)		100.0 %	0.29 [-0.04, 0.62]
Subtotal (95% CI)	71		72		-	100.0 %	0.29 [-0.04, 0.62]
Heterogeneity: not applicable							
Test for overall effect: Z = 1.71 (P	= 0.087)						
3 Calcipotriol (12 wks) vs. combin	ed calcipotric	l + BMD (4 wks); tl	hen calcipotriol	(w/dy) % comb	ined calcipotriol + BMD ((w/e) (8 wks)	
Kragballe 2004	327 -	0.659 (0.331)	323	-0.69 (0.285)		100.0 %	0.10 [-0.05, 0.25]
Subtotal (95% CI)	327		323		•	100.0 %	0.10 [-0.05, 0.25]
Heterogeneity: not applicable	527		0-0			10000 /0	0120 [0109, 0129]
Test for overall effect: $Z = 1.28$ (P	= 0.20)						
4 Calcipotriol (6 wks) vs. clobetas	/	(2 wks); then calcip	otriol (4 wks)				
Subtotal (95% CI)	0) O				Not estimable
Heterogeneity: not applicable							
Test for overall effect: not applicab	le						
5 Calcipotriol (6 wks) vs. calcipotr	iol OM, fluoc	inonide acetonide C	DN (2 wks); the	n calcipotriol tw	vice daily (4 wks)		
Wozel 2001	19	6.5 (3.86)	19	4.27 (2.64)		→ I00.0 %	0.66 [0.01, 1.32]
Subtotal (95% CI)	19		19			- 100.0 %	0.66 [0.01, 1.32]
Heterogeneity: not applicable							
Test for overall effect: $Z = 1.98$ (P	= 0.048)						
6 Calcipotriol (6 wks) vs. halometa	asone OM, ca	lcipotriol ON (2 wk	s); then calcipot	riol twice daily ((w/dy), halometasone (w/e	e) (2 wks); then	calcipotriol twice daily (2
wks)							
Yang 2009	36	4.5 (2.5)	40	2.3 (1.2)		→ I00.0 %	1.13 [0.64, 1.62]
Subtotal (95% CI)	36		40		_	100.0 %	1.13 [0.64, 1.62]
Heterogeneity: not applicable							
Test for overall effect: $Z = 4.55$ (P	< 0.00001)						
7 Calcipotriol ON, clobetasol pro	pionate OM (2 to 4 wks); then c	alcipotriol twice	e daily (to wk 12	!) vs. calcitriol ON, clobet	asol propionate	e OM (2 to 4 wks); then
calcitriol twice daily (to wk 12)							
						1	
				-	-0.5 0 0.5	I	
				Favours vitamin I	D regimen Favours co	mplex regimen	
							(Continued)

Study or subgroup	Vitamin D regimen	Com	plex regimen		Std. Mean Difference	Weight	(Continued Sta Mea Differenc
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% (
Lahfa 2003	64	-0.82 (0.3)	61	-0.74 (0.3)		100.0 %	-0.27 [-0.62, 0.09
Subtotal (95% CI)	64		61			100.0 %	-0.27 [-0.62, 0.09
Heterogeneity: not applic	able						
Test for overall effect: Z =	= 1.47 (P = 0.14)						
8 Combined calcipotriol	+ BMD (4 wks); then	placebo ointment tw	ice daily (8 wk	s) vs. combined	calcipotriol + BMD (4 wl	<s); calcipo<="" td="" then=""><td>triol ointment twice daily</td></s);>	triol ointment twice daily
(8 wks) White 2006 (P)	376	-0.33 (0.48)	383	-0.45 (0.49)	-	100.0 %	0.25 [0.10, 0.39
Subtotal (95% CI)	376		383		•	100.0 %	0.25 [0.10, 0.39
Heterogeneity: not applic	able						
Test for overall effect: Z =	= 3.39 (P = 0.00070)						
9 Combined calcipotriol	+ BMD (4 wks); then	placebo ointment tw	rice daily (8 wk	s) vs. combined	calcipotriol + BMD (4 w	ks); then calcipo	triol (w/dy) + combined
calcipotriol + BMD (w/	, , ,			0.50 (6.55)			
White 2006 (P)	376	-0.33 (0.48)	377	-0.58 (0.35)		100.0 %	0.59 [0.45, 0.74
Subtotal (95% CI)	376		377		•	100.0 %	0.59 [0.45, 0.74
Heterogeneity: not applic Test for overall effect: Z =							
				(0, 1,)			
10 Combined calcipotrio combined calcipotriol	, ,		nt twice daily	(8 wks) vs. com	bined calcipotriol + BIML) (4 wks); then	calcipotriol (w/dy) +
White 2006 (H)	383 383	-0.45 (0.49)	377	-0.58 (0.35)	-	100.0 %	0.30 [0.16, 0.45
Subtotal (95% CI)	383		377		•	100.0 %	0.30 [0.16, 0.45
Heterogeneity: not applic Test for overall effect: Z =							
	,						
II Combined calcipotrio	I + BMD (8 wks); the	n calcipotriol (4 wks)	vs. combined	calcipotriol + BI	MD (4 wks); then calcipo	triol (w/dy) % c	ombined calcipotriol +
BMD (w/e) (8 wks) Kragballe 2004	322 -	0.646 (0.317)	323	-0.69 (0.285)	-	100.0 %	0.15 [-0.01, 0.30
Subtotal (95% CI)	322	~ /	323	()	•	100 0 %	0.15 [-0.01, 0.30
Heterogeneity: not applic			525			100.0 /0	0.19 [0.01, 0.90
Test for overall effect: Z =							
			1				
12 Tacalcitol (8 wks) vs. c Ortonne 2004	ombined calcipotriol - 252	-0.38 (0.46)		-0.59 (0.39)		100.0 %	0.49 [0.31, 0.67
		0.00 (01.0)		0.07 (0.07)			-
Subtotal (95% CI)	252		249		-	100.0 %	0.49 [0.31, 0.67
Heterogeneity: not applic							
Test for overall effect: Z = Test for subgroup differer	,	-10(P-0.00) 12 -	.86%				
iest ior subgroup differen	ices. CIII - 70.21, dī	- 10 (1 - 0.00), 1 ² -	.00/0				
				-	-0.5 0 0.5		

Analysis 13.4. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

Outcome: 4 PAGI

Study or subgroup 🛛 🕔	/itamin D regimen N	C Mean(SD)	Complex regimen N	Mean(SD)	S Me Differer IV,Random,9	ice W	eight	Std. Mean Difference IV,Random,95% CI
Calcipotriol (12 wks) vs. o	combined calcipotri	ol + BMD (8 wł	s); then calcipotri	ol (4 wks)				
Kragballe 2004	283	-3.52 (0.97)	294	-3.37 (1.18)		100).0 %	-0.14 [-0.30, 0.02]
Subtotal (95% CI)	283		294		•	100.)% -0.1 4	£ [-0.30, 0.02]
Heterogeneity: not applicat	ble							
Test for overall effect: $Z =$	I.66 (P = 0.097)							
2 Calcipotriol (12 wks) vs. o Subtotal (95% CI)	combined calcipotri 0	ol + BMD (4 wł	s); then calcipotri 0	ol (8 wks)				Not estimable
Heterogeneity: not applicab	-		Ŭ					
Test for overall effect: not a								
3 Calcipotriol (12 wks) vs. o	• •	ol + BMD (4 wł	s); then calcipotri	ol (w/dy) % com	bined calcipotrio	+ BMD (w/e) (8	wks)	
Kragballe 2004		-3.52 (0.97)	, ,	-3.62 (0.98)).0 %	0.10 [-0.06, 0.26]
Subtotal (95% CI)	283		302		•	100.)% 0.1(0 [-0.06, 0.26]
Heterogeneity: not applicab	ble							
Test for overall effect: Z =	I.24 (P = 0.22)							
4 Calcipotriol (6 wks) vs. cl	obetasol propionat	e (2 wks); then a	alcipotriol (4 wks))				
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicat	ble							
Test for overall effect: not a	pplicable							
5 Calcipotriol (6 wks) vs. ca	alcipotriol OM, fluo	cinonide acetoni	de ON (2 wks); tl	nen calcipotriol t	wice daily (4 wks)		
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicab	ble							
Test for overall effect: not a	• •							
6 Calcipotriol (6 wks) vs. ha	alometasone OM, c	alcipotriol ON (3	2 wks); then calcip	otriol twice daily	/ (w/dy), halometi	asone (w/e) (2 wk	s); then calci	potriol twice daily (2
wks)	0		0					Net estimable
Subtotal (95% CI)	-		0					Not estimable
Heterogeneity: not applicab Test for overall effect: not a								
7 Calcipotriol ON, clobetas		(2 to 4 w/s), th	en calcinotriol twi	ce daily (to wk	12) vs. calcitriol C	N clobetasol pro	nionate OM	(2 to 4 w/s); then
calcitriol twice daily (to w		(2 to 1 Wio), th			(2) V3. Calcia (0) C	int, clobetasor pro		
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicab	ble							
Test for overall effect: not a	pplicable							
8 Combined calcipotriol +	BMD (4 wks); then	placebo ointme	nt twice daily (8 v	vks) vs. combine	d calcipotrio + B	MD (4 wks); then	calcipotriol	ointment twice daily
(8 wks)								
				I				
				-	-0.5 0	0.5 I		
				Favours vitamin	D regimen F	avours complex regi	nen	
								(Continued

(Continued . . .)

Study or subgroup Vita	min D regimen	M (CD)	Complex regimen	M (CD)	Std. Mean Difference	Weight	(Continued Std. Mean Difference
\\/\-;+- 2007 (D)	N	Mean(SD)	N	Mean(SD)	IV,Random,95% CI	100.0 %	IV,Random,95% CI
White 2006 (P)	376	-2.25 (1.79)	383	-2.73 (1.67)		100.0 %	0.28 [0.13, 0.42]
Subtotal (95% CI)	376		383		*	100.0 %	0.28 [0.13, 0.42]
Heterogeneity: not applicable							
Test for overall effect: $Z = 3.8$	` /						
9 Combined calcipotriol + B№		i placebo ointr	nent twice daily (8 v	vks) vs. combine	d calcipotrio + BMD (4 v	wks); then calcip	otriol (w/dy) + combined
calcipotriol + BMD (w/e) (8 White 2006 (P)	,	-2.25 (1.79)	377	-3.44 (1.57)	-	100.0 %	0.71 [0.56, 0.85]
Subtotal (95% CI)	376		377		•	100.0 %	0.71 [0.56, 0.85]
Heterogeneity: not applicable							
Test for overall effect: $Z = 9.4$	0 (P < 0.00001)						
10 Combined calcipotriol + B combined calcipotriol + BN White 2006 (H)	1D (w/e) (8 wks			y (8 wks) vs. cor -3.44 (1.57)	nbined calcipotriol + BM	D (4 wks); ther 100.0 %	0.44 [0.29, 0.58 ⁻
Subtotal (95% CI)	383	. ,	377	. ,	•	100.0 %	0.44 [0.29, 0.58]
Heterogeneity: not applicable	<i>J</i> 0 <i>J</i>		3//			100.0 70	0.44 [0.29, 0.98]
Test for overall effect: $Z = 5.9$	(P < 0.00001)						
Test for overall effect. $\Sigma = 5.7$	5 (1 < 0.00001)						
II Combined calcipotriol + B	MD (8 wks); the	en calcipotriol	(4 wks) vs. combine	d calcipotriol +	BMD (4 wks); then calcip	otriol (w/dy) %	combined calcipotriol +
BMD (w/e) (8 wks)	20.4		202	2 (2 (0 0 0)		100.0 %	
Kragballe 2004	294	-3.37 (1.18)	302	-3.62 (0.98)		100.0 %	0.23 [0.07, 0.39]
Subtotal (95% CI)	294		302		•	100.0 %	0.23 [0.07, 0.39]
Heterogeneity: not applicable							
Test for overall effect: $Z = 2.80$	0 (P = 0.0050)						
12 Tacalcitol (8 wks) vs. comb	ined calcipotriol	+ BMD (4 wk	s); then calcipotriol	(4 wks)			
Ortonne 2004	247	-2.4 (1.38)	246	-3.15 (1.39)		100.0 %	0.54 [0.36, 0.72]
Subtotal (95% CI)	247		246		•	100.0 %	0.54 [0.36, 0.72]
Heterogeneity: not applicable	= -/		210				
Test for overall effect: $Z = 5.9$	0 (P < 0.00001)						
Test for subgroup differences:	,	F = 6 (P = 0.00)), I ² =92%				
5							
				-	-0.5 0 0.5	I	
				Favours vitamin I		mplex regimen	

Analysis 13.5. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D regimen N	Mean(SD)	Complex regimen N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
Calcipotriol (12 wks) v	vs. combined calcipotri	ol + BMD (8	wks); then calcipotri	ol (4 wks)			
Kragballe 2004	283	1.62 (0.86)	294	1.73 (0.92)		100.0 %	-0.12 [-0.29, 0.04]
Subtotal (95% CI) Heterogeneity: not appli Test for overall effect: Z	cable		294		•	100.0 %	-0.12 [-0.29, 0.04]
2 Calcipotriol (12 wks) v	vs. combined calcipotr	ol + BMD (4	wks); then calcipotri	ol (8 wks)	_		
Saraceno 2007	71	3.04 (3.76)	72	2.11 (2.56)		100.0 %	0.29 [-0.04, 0.62]
Subtotal (95% CI) Heterogeneity: not appli Test for overall effect: Z	cable		72		•	100.0 %	0.29 [-0.04, 0.62]
,				,	bined calcipotriol + BMD	. , . ,	
Kragballe 2004	283	1.62 (0.86)	302	1.51 (0.89)		100.0 %	0.13 [-0.04, 0.29]
Subtotal (95% CI)) 283		302		•	100.0 %	0.13 [-0.04, 0.29]
Heterogeneity: not appli Test for overall effect: Z 4 Calcipotriol (6 wks) vs	= 1.51 (P = 0.13)	e (2 wks); the	n calcipotriol (4 wks)	1			
Austad 1998	46	-3.17 (0.82)	46	-3.67 (0.84)		100.0 %	0.60 [0.18, 1.02
Subtotal (95% CI)) 46		46		•	100.0 %	0.60 [0.18, 1.02]
Heterogeneity: not appli Test for overall effect: Z 5 Calcipotriol (6 wks) vs	= 2.80 (P = 0.0051) calcipotriol OM, fluo				wice daily (4 wks)		
Wozel 2001	19	6.5 (3.86)	19	4.27 (2.64)		100.0 %	0.66 [0.01, 1.32
Subtotal (95% CI) Heterogeneity: not appli Test for overall effect: Z	cable = 1.98 (P = 0.048)		19		•	1 00.0 %	0.66 [0.01, 1.32]
	. halometasone OM, c	alcipotriol ON	I (2 wks); then calcip	otriol twice daily	/ (w/dy), halometasone (w	/e) (2 wks); the	n calcipotriol twice daily (2
wks) Yang 2009	36	-1.86 (0.93)	40	-2.23 (0.86)		100.0 %	0.41 [-0.05, 0.86
Subtotal (95% CI)			40		•	100.0 %	0.41 [-0.05, 0.86
Heterogeneity: not appli Test for overall effect: Z 7 Calcipotriol ON, clobe calcitriol twice daily (to	= 1.76 (P = 0.078) etasol propionate OM	(2 to 4 wks);	then calcipotriol twi	ce daily (to wk I	2) vs. calcitriol ON, clobe	etasol propiona	te OM (2 to 4 wks); then
				-2 Favours vitamin	-1 0 1 D regimen Favours co	2 mplex regimen	(Continued

					Std.		(Continue Sto
Study or subgroup	Vitamin D regimen	Comp	lex regimen		Mean Difference	Weight	Mea Differenc
Study of Subgroup	N	Mean(SD)	N	Mean(SD)	IV,Random,95% Cl	* * Cigiric	IV,Random,95% C
Lahfa 2003	64	-4.69 (1.3)	61	-4.44 (1.28)		100.0 %	-0.19 [-0.54, 0.16
Subtotal (95% CI)	64	~ /	61	· · /	-	100.0 %	-0.19 [-0.54, 0.16
Heterogeneity: not applic Test for overall effect: Z =	able		01			100.0 /0	-0.17 [-0.94, 0.10
8 Combined calcipotriol	+ BMD (4 wks); then	placebo ointment tw	ice daily (8 w	rks) vs. combined	d calcipotriol + BMD (4 v	vks); then calcip	otriol ointment twice dail
(8 wks) White 2006 (P)	376	2.49 (0.99)	383	2.23 (0.96)		100.0 %	0.27 [0.12, 0.41
Subtotal (95% CI)	376		383		•	100.0 %	0.27 [0.12, 0.41
Heterogeneity: not applic	able						
Test for overall effect: Z =	= 3.65 (P = 0.00026)						
9 Combined calcipotriol	+ BMD (4 wks); then	placebo ointment tw	vice daily (8 v	vks) vs. combine	d calcipotrio + BMD (4 v	wks); then calcip	ootriol (w/dy)+ combined
calcipotriol + BMD (w/ White 2006 (P)	'e) (8 wks) 376	2.49 (0.99)	377	1.98 (0.99)		100.0 %	0.51 [0.37, 0.66
Subtotal (95% CI)	376		377		•	100.0 %	0.51 [0.37, 0.66
Heterogeneity: not applic	able						
Test for overall effect: Z =	= 6.95 (P < 0.00001)						
10 Combined calcipotrio	I + BMD (4 wks); the	en calcipotriol ointme	nt twice dail	/ (8 wks) vs. con	nbined calcipotriol + BM	D (4 wks); ther	n calcipotriol (w/dy) +
combined calcipotriol	+ BMD (w/e) (8 wks)		. ,			
White 2006 (H)	383	2.23 (0.96)	377	1.98 (0.99)	-	100.0 %	0.26 [0.11, 0.40
Subtotal (95% CI)	383		377		•	100.0 %	0.26 [0.11, 0.40
Heterogeneity: not applic	able						
Test for overall effect: Z =	= 3.52 (P = 0.00044)						
II Combined calcipotrio	I + BMD (8 wks); the	n calcipotriol (4 wks)	vs. combine	d calcipotriol + E	3MD (4 wks); then calcip	otriol (w/dy) %	combined calcipotriol +
BMD (w/e) (8 wks)				1.5.1 (0.00)		100.0.0/	
Kragballe 2004	294	1.73 (0.92)	302	1.51 (0.89)		100.0 %	0.24 [0.08, 0.40
Subtotal (95% CI)	294		302		•	100.0 %	0.24 [0.08, 0.40
Heterogeneity: not applic							
Test for overall effect: Z =	– 2.95 (P – 0.0032)						
12 Tacalcitol (8 wks) vs. o	combined calcipotriol	+ BMD (4 wks); then	calcipotriol ((4 wks)			
Ortonne 2004	247	-2.45 (1.31)	246	-3.16 (1.33)		100.0 %	0.54 [0.36, 0.72
Subtotal (95% CI)	247		246		•	100.0 %	0.54 [0.36, 0.72
Heterogeneity: not applic							
Test for overall effect: Z =	= 5.86 (P < 0.00001)						
Test for subgroup differer	nces: Chi ² = 55.51, df	$= (P = 0.00), ^2 =$	80%				
				1			
				-2	-1 0 1	2	

Analysis 13.6. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

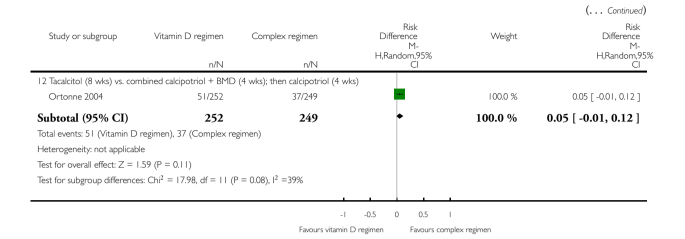
Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D regimen	Complex regimen	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Calcipotriol (12 wks) vs. c	combined calcipotriol + BM	1D (8 wks); then calcipotriol ((4 wks)		
Kragballe 2004	47/327	30/322	-	100.0 %	0.05 [0.00, 0.10]
Subtotal (95% CI)	327	322	•	100.0 %	0.05 [0.00, 0.10]
Total events: 47 (Vitamin D	regimen), 30 (Complex re	gimen)			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 2$	2.00 (P = 0.045)				
,	combined calcipotriol + BM	1D (4 wks); then calcipotriol ((8 wks)		
Saraceno 2007	17/75	14/75		100.0 %	0.04 [-0.09, 0.17]
Subtotal (95% CI)	75	75	+	100.0 %	0.04 [-0.09, 0.17]
Total events: 17 (Vitamin D	regimen), 14 (Complex re	gimen)			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 0$	0.61 (P = 0.54)				
,			(w/dy) % combined calcipotric	. , , ,	rs)
Kragballe 2004	47/327	21/322		100.0 %	0.08 [0.03, 0.13]
Subtotal (95% CI)	327	322	•	100.0 %	0.08 [0.03, 0.13]
Total events: 47 (Vitamin D	regimen), 21 (Complex re	gimen)			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 3$	8.30 (P = 0.00096)				
4 Calcipotriol (6 wks) vs. clo		, , , , ,			
Austad 1998	3/49	3/49	-	100.0 %	0.0 [-0.09, 0.09]
Subtotal (95% CI)	49	49	+	100.0 %	0.0 [-0.09, 0.09]
Total events: 3 (Vitamin D r	regimen), 3 (Complex regin	nen)			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
		()	n calcipotriol twice daily (4 wks	,	
Wozel 2001	0/19	0/19	-	100.0 %	0.0 [-0.10, 0.10]
Subtotal (95% CI)	19	19	+	100.0 %	0.0 [-0.10, 0.10]
Total events: 0 (Vitamin D r	regimen), 0 (Complex regin	nen)			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
6 Calcipotriol (6 wks) vs. ha	lometasone OM, calcipotri	ol ON (2 wks); then calcipotr	riol twice daily (w/dy), halomet	asone (w/e) (2 wks);	then calcipotriol twice daily (2
wks)					
		-	-0.5 0 0.5 I		
		Eavours vitami	in D regimen Favours comp	lex regimen	

Study or subgroup	Vitamin D regimen	Complex regimen	Risk Difference	Weight	Risk Difference
	n/N	n/N	M- H,Random,95% Cl		M- H,Random,S Cl
Yang 2009	0/36	0/40	+	100.0 %	0.0 [-0.05, 0.05]
Subtotal (95% CI)	36	40	•	100.0 %	0.0 [-0.05, 0.05]
Fotal events: 0 (Vitamin D n Heterogeneity: not applicab Fest for overall effect: Z = 0	egimen), 0 (Complex regin le 1.0 (P = 1.0)	nen)			
		wks); then calcipotriol twice	daily (to wk 12) vs. calcitriol	ON, clobetasol propio	onate OM (2 to 4 wks); then
calcitriol twice daily (to w Lahfa 2003	k 12) 8/64	8/61	-	100.0 %	-0.0 [-0.12, 0.11]
Subtotal (95% CI)	64	61	+	100.0 %	-0.01 [-0.12, 0.11]
Total events: 8 (Vitamin D n	egimen), 8 (Complex regin	nen)			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = C$	0.10 (P = 0.92)				
3 Combined calcipotriol + 8	3MD (4 wks); then placebo	o ointment twice daily (8 wks)) vs. combined calcipotriol +	BMD (4 wks); then ca	lcipotriol ointment twice daily
(8 wks) White 2006 (P)	79/376	48/383	-	100.0 %	0.08 [0.03, 0.14]
Subtotal (95% CI)	376	383	•	100.0 %	0.08 [0.03, 0.14]
Total events: 79 (Vitamin D		gimen)			
Heterogeneity: not applicab					
Test for overall effect: $Z = 3$	· · · ·				
		o ointment twice daily (8 wks) vs. combined calcipotriol +	BMD (4 wks); then ca	alcipotriol (w/dy) + combined
calcipotriol + BMD (w/e) White 2006 (P)	(8 wks) 79/376	36/377		100.0 %	0.11 [0.06, 0.17]
Subtotal (95% CI)	376	377	•	100.0 %	0.11 [0.06, 0.17]
Fotal events: 79 (Vitamin D	regimen), 36 (Complex re	gimen)			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 4$.43 (P < 0.00001)				
0 Combined calcinotriol +	BMD (4 wks): then calcin	otriol ointment twice daily (8	8 wks) vs. combined calcinot	riol + BMD (4 wks): 1	then calcinotrial (w/dv) +
combined calcipotriol +	. , , ,				anen calcipotnor (widy)
White 2006 (H)	48/383	36/377	-	100.0 %	0.03 [-0.01, 0.07]
Subtotal (95% CI)	383	377	•	100.0 %	0.03 [-0.01, 0.07]
Fotal events: 48 (Vitamin D	regimen), 36 (Complex re	gimen)			
Heterogeneity: not applicab	le				
Test for overall effect: $Z = I$.31 (P = 0.19)				
L Combined calcinotrial +	BMD (8 w/s); then calcin	otriol (4 wks) vs. combined c	alcipotrial + BMD (4 w/s); t	hen calcinotrial (w/dv) % combined calcinotrial +
BMD (w/e) (8 wks)	brib (0 wis), then earlip				
Kragballe 2004	30/322	21/322	+	100.0 %	0.03 [-0.01, 0.07
Subtotal (95% CI)	322	322	•	100.0 %	0.03 [-0.01, 0.07]
Total events: 30 (Vitamin D				100.0 %	0.03 [-0.01, 0.07
Heterogeneity: not applicab		5 ⁽¹⁾ (1)			
Test for overall effect: $Z = 1$					
z = 1					
				-	
		-	I -0.5 0 0.5	I	

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Analysis 13.7. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D regimen	Complex regimen	Risk Difference M-	Weight	Risk Difference M-
	n/N	N n/N	H,Random,95% Cl		H,Random,95% Cl_
Calcipotriol (12 wks) vs.	combined calcipotriol + BM	D (8 wks); then calcipotriol	(4 wks)		
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D	regimen), 0 (Complex regim	nen)			
Heterogeneity: not applica	able				
Test for overall effect: not	applicable				
2 Calcipotriol (12 wks) vs.	combined calcipotriol + BM	D (4 wks); then calcipotriol	(8 wks)		
Saraceno 2007	0/75	3/75	-	100.0 %	-0.04 [-0.09, 0.01]
Subtotal (95% CI)	75	75	•	100.0 %	-0.04 [-0.09, 0.01]
Total events: 0 (Vitamin D	regimen), 3 (Complex regim	nen)			
Heterogeneity: not applica	able				
Test for overall effect: Z =	1.55 (P = 0.12)				
			-1 -0.5 0 0.5	I	
		Favours vitan	nin D regimen Favours co	omplex regimen	
					(Continued)

Study or subgroup	Vitamin D regimen	Complex regimen	Difference M-	Weight	Difference
	n/N	n/N	H,Random,95% Cl		H,Random C
,			w/dy) % combined calcipotric	ol + BMD (w/e) (8 w	,
Subtotal (95% CI)	0	0			Not estimable
	regimen), 0 (Complex regin	nen)			
Heterogeneity: not applicat					
Test for overall effect: not a 4 Calcipotriol (6 w/s) vs. cl	pplicable obetasol propionate (2 wks) then calcingtrial (4 who)			
Austad 1998	0/49	0/49	-	100.0 %	0.0 [-0.04, 0.04
Subtotal (95% CI)	49	49	•	100.0 %	0.0 [-0.04, 0.04
(-	regimen), 0 (Complex regin			10000 /0	
Heterogeneity: not applicat		,			
Test for overall effect: $Z = 0$					
	. ,	acetonide ON (2 wks); then	a calcipotriol twice daily (4 wks	s)	
Wozel 2001	0/19	0/19	#	100.0 %	0.0 [-0.10, 0.10
Subtotal (95% CI)	19	19	+	100.0 %	0.0 [-0.10, 0.10
	regimen), 0 (Complex regin				,
Heterogeneity: not applicat	8 , (1 8				
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
	alometasone OM, calcipotrio	ol ON (2 wks); then calcipotr	iol twice daily (w/dy), halomet	tasone (w/e) (2 wks);	then calcipotriol twice daily (
wks) Yang 2009	0/36	0/40	•	100.0 %	0.0 [-0.05, 0.05
Subtotal (95% CI)	36	40	•	100.0 %	0.0 [-0.05, 0.05
	regimen), 0 (Complex regin				
Heterogeneity: not applicat		,			
Test for overall effect: $Z = 0$					
7 Calcipotriol ON, clobetas	sol propionate OM (2 to 4	wks); then calcipotriol twice	daily (to wk 12) vs. calcitriol (ON, clobetasol propie	onate OM (2 to 4 wks); then
calcitriol twice daily (to w	,				
Lahfa 2003	0/64	1/61		100.0 %	-0.02 [-0.06, 0.03
Subtotal (95% CI)	64	61	+	100.0 %	-0.02 [-0.06, 0.03
Total events: 0 (Vitamin D r	regimen), I (Complex regin	nen)			
Heterogeneity: not applicat					
Test for overall effect: $Z = 0$. ,				
	BMD (4 wks); then placebo	o ointment twice daily (8 wks)) vs. combined calcipotriol + E	3MD (4 wks); then ca	lcipotriol ointment twice dail;
(8 wks) White 2006 (P)	7/376	5/383		100.0 %	0.01 [-0.01, 0.02
Subtotal (95% CI)	376	383	•	100.0 %	0.01 [-0.01, 0.02
	regimen), 5 (Complex regin				
Heterogeneity: not applicat	0 / ()	,			
Test for overall effect: $Z = 0$					
9 Combined calcipotriol +	BMD (4 wks); then placebo	o ointment twice daily (8 wks)) vs. combined calcipotriol + I	BMD (4 wks); then ca	alcipotriol (w/dy) + combined
calcipotriol + BMD (w/e)	, ,				
White 2006 (P)	7/376	8/377	Ţ	100.0 %	0.00 [-0.02, 0.02
Subtotal (95% CI)	376	377		100.0 %	0.00 [-0.02, 0.02
-					
		-	I -0.5 0 0.5 I		

Study or subgroup	Vitamin D regimen	Complex regimen	Risk Difference M- H,Random,95%	Weight	(Continued) Risk Difference M- H.Random,95
	n/N	n/N	CI		Cl
Total events: 7 (Vitamin D reg	gimen), 8 (Complex regin	nen)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.2$	26 (P = 0.80)				
10 Combined calcipotriol + I	BMD (4 wks); then calcip	ootriol ointment twice daily (8	3 wks) vs. combined calcipot	riol + BMD (4 wks); t	hen calcipotriol (w/dy) +
combined calcipotriol + B	MD (w/e) (8 wks)				
White 2006 (H)	5/383	8/377	-	100.0 %	-0.01 [-0.03, 0.01]
Subtotal (95% CI)	383	377	•	100.0 %	-0.01 [-0.03, 0.01]
Total events: 5 (Vitamin D reg	gimen), 8 (Complex regin	nen)			
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.8$	87 (P = 0.39)				
Combined calcipotriol + E	BMD (8 wks): then calcip	otriol (4 wks) vs. combined c	alcipotriol + BMD (4 wks); t	hen calcipotriol (w/dv) % combined calcipotriol +
II Combined calcipotriol + E BMD (w/e) (8 wks)	BMD (8 wks); then calcip	otriol (4 wks) vs. combined c	alcipotriol + BMD (4 wks); tl	hen calcipotriol (w/dy)) % combined calcipotriol +
II Combined calcipotriol + E BMD (w/e) (8 wks) Subtotal (95% CI)	BMD (8 wks); then calcip	otriol (4 wks) vs. combined c 0	alcipotriol + BMD (4 wks); tl	hen calcipotriol (w/dy)) % combined calcipotriol + Not estimable
BMD (w/e) (8 wks)	0	0	alcipotriol + BMD (4 wks); tl	hen calcipotriol (w/dy	
BMD (w/e) (8 wks) Subtotal (95% CI)	0 gimen), 0 (Complex regin	0	alcipotriol + BMD (4 wks); tl	hen calcipotriol (w/dy	
BMD (w/e) (8 wks) Subtotal (95% CI) Total events: 0 (Vitamin D reg	0 gimen), 0 (Complex regin	0	alcipotriol + BMD (4 wks); tl	hen calcipotriol (w/dy	
BMD (w/e) (8 wks) Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app	0 gimen), 0 (Complex regin plicable	0 nen)		hen calcipotriol (w/dy	
BMD (w/e) (8 wks) Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable	0 gimen), 0 (Complex regin plicable	0 nen)		hen calcipotriol (w/dy	
BMD (w/e) (8 wks) Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app 12 Tacalcitol (8 wks) vs. comb Ortonne 2004	0 gimen), 0 (Complex regin plicable bined calcipotriol + BMD	0 nen) (4 wks); then calcipotriol (4 v		100.0 %	Not estimable 0.02 [-0.01, 0.05]
BMD (w/e) (8 wks) Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app 12 Tacalcitol (8 wks) vs. comb Ortonne 2004 Subtotal (95% CI)	0 gimen), 0 (Complex regin plicable bined calcipotriol + BMD I 1/252 252	0 nen) (4 wks); then calcipotriol (4 v 6/249 249			Not estimable
BMD (w/e) (8 wks) Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app 12 Tacalcitol (8 wks) vs. comb Ortonne 2004	0 gimen), 0 (Complex regin plicable bined calcipotriol + BMD 11/252 252 egimen), 6 (Complex reg	0 nen) (4 wks); then calcipotriol (4 v 6/249 249		100.0 %	Not estimable 0.02 [-0.01, 0.05]
BMD (w/e) (8 wks) Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app 12 Tacalcitol (8 wks) vs. comb Ortonne 2004 Subtotal (95% CI) Total events: 11 (Vitamin D reg	0 gimen), 0 (Complex regin plicable bined calcipotriol + BMD I 1/252 252 egimen), 6 (Complex reg	0 nen) (4 wks); then calcipotriol (4 v 6/249 249		100.0 %	Not estimable 0.02 [-0.01, 0.05]
BMD (w/e) (8 wks) Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app 12 Tacalcitol (8 wks) vs. comb Ortonne 2004 Subtotal (95% CI) Total events: 11 (Vitamin D re Heterogeneity: not applicable	0 gimen), 0 (Complex reginer plicable bined calcipotriol + BMD I 1/252 252 egimen), 6 (Complex reg	0 nen) (4 wks); then calcipotriol (4 w 6/249 249 imen)		100.0 %	Not estimable 0.02 [-0.01, 0.05]

Favours vitamin D regimen Favours complex regimen

Analysis 13.8. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

Outcome: 8 Withdrawals due to treatment failure

Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app	0 imen), 0 (Complex regin licable	n/N 1D (8 wks); then calcipotriol (4 0 nen) 1D (4 wks); then calcipotriol (8 4/75			M- H,Random, CI Not estimable
Subtotal (95% CI) Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app	0 imen), 0 (Complex regin licable nbined calcipotriol + BM	0 nen) 1D (4 wks); then calcipotriol (8			Not estimable
Total events: 0 (Vitamin D reg Heterogeneity: not applicable Test for overall effect: not app	imen), 0 (Complex regin licable nbined calcipotriol + BM	nen) 1D (4 wks); then calcipotriol (8	3 wks)		Not estimable
Heterogeneity: not applicable Test for overall effect: not app	licable nbined calcipotriol + BM	1D (4 wks); then calcipotriol (8	3 wks)		
Test for overall effect: not app	licable nbined calcipotriol + BM	. ,	3 wks)		
	mbined calcipotriol + BM	. ,	3 wks)		
2 Calcipotriol (12 wks) vs. cor		. ,	3 wks)		
	20/75	4/75			
Saraceno 2007				100.0 %	0.21 [0.10, 0.33]
Subtotal (95% CI)	75	75	•	100.0 %	0.21 [0.10, 0.33]
Total events: 20 (Vitamin D re	gimen), 4 (Complex regi	men)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 3.7$	2 (P = 0.00020)				
3 Calcipotriol (12 wks) vs. cor	mbined calcipotriol + BM	1D (4 wks); then calcipotriol (w	w/dy) % combined calcipotrio	I + BMD (w/e) (8 wks))
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D reg	imen), 0 (Complex regin	nen)			
Heterogeneity: not applicable					
Test for overall effect: not app	licable				
4 Calcipotriol (6 wks) vs. clob					
Austad 1998	0/49	0/49		100.0 %	0.0 [-0.04, 0.04]
Subtotal (95% CI)	49	49	•	100.0 %	0.0 [-0.04, 0.04]
Total events: 0 (Vitamin D reg	imen), 0 (Complex regin	nen)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
5 Calcipotriol (6 wks) vs. calcij	potriol OM, fluocinonide	e acetonide ON (2 wks); then o	calcipotriol twice daily (4 wks)	
Wozel 2001	0/19	0/19		100.0 %	0.0 [-0.10, 0.10]
Subtotal (95% CI)	19	19	+	100.0 %	0.0 [-0.10, 0.10]
Total events: 0 (Vitamin D reg	imen), 0 (Complex regin	nen)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
6 Calcipotriol (6 wks) vs. halor	metasone OM, calcipotri	ol ON (2 wks); then calcipotric	ol twice daily (w/dy), halomet	asone (w/e) (2 wks); th	en calcipotriol twice daily (2
wks)	0.57	0/40		100.0.57	
Yang 2009	0/36	0/40	-	100.0 %	0.0 [-0.05, 0.05]
Subtotal (95% CI)	36	40	•	100.0 %	0.0 [-0.05, 0.05]
			I -0.5 0 0.5 I		
		Favours vitamir			
				0	(Continued

(... Continued)

Study or subgroup	Vitamin D regimen	Complex regimen	Difference M-	Weight	Difference M
	n/N	n/N	H,Random,95% Cl		H,Random, C
Total events: 0 (Vitamin D r	egimen), 0 (Complex regim	nen)			
Heterogeneity: not applicab	e				
Test for overall effect: $Z = 0$.0 (P = 1.0)				
' Calcipotriol ON, clobetas	ol propionate OM (2 to 4	wks); then calcipotriol twice	daily (to wk 12) vs. calcitriol (DN, clobetasol propion	ate OM (2 to 4 wks); then
calcitriol twice daily (to w Lahfa 2003	< 2) 4/64	2/61		100.0 %	0.03 [-0.04, 0.10
	()	(1		100.0.0/	
Subtotal (95% CI)	64	61	-	100.0 %	0.03 [-0.04, 0.10
	egimen), 2 (Complex regim	nen)			
leterogeneity: not applicab					
test for overall effect: $Z = 0$	· /				
	3MD (4 wks); then placebo	ointment twice daily (8 wks)) vs. combined calcipotriol + E	3MD (4 wks); then calc	ipotriol ointment twice dail
(8 wks)	~	2			
Subtotal (95% CI)	0	0			Not estimable
	egimen), 0 (Complex regim	nen)			
leterogeneity: not applicab	e				
est for overall effect: not a	oplicable				
Combined calcipotriol +	3MD (4 wks); then placebo	ointment twice daily (8 wks)) vs. combined calcipotriol + I	3MD (4 wks); then calc	ipotriol (w/dy) + combined
calcipotriol + BMD (w/e)	(8 wks) 0	0			Not estimable
	egimen), 0 (Complex regim				100000000000000000000000000000000000000
Heterogeneity: not applicab					
est for overall effect: not a					
est for overall cheet. Not a	plicable				
0 Combined calcipotriol +	BMD (4 wks); then calcip	otriol ointment twice daily (8 wks) vs. combined calcipotr	iol + BMD (4 wks); th	en calcipotriol (w/dy)+
combined calcipotriol + Subtotal (95% CI)	BMD (w/e) (8 wks) 0	0			Not estimabl
	egimen), 0 (Complex regim	nen)			
leterogeneity: not applicab					
est for overall effect: not a					
	BMD (8 wks); then calcipo	otriol (4 wks) vs. combined c	alcipotriol + BIMD (4 wks); th	en calcipotriol (w/dy) 9	% combined calcipotriol +
BMD (w/e) (8 wks)					
Subtotal (95% CI)	0	0			Not estimable
	egimen), 0 (Complex regim	nen)			
Heterogeneity: not applicab					
est for overall effect: not a	oplicable				
2 Tacalcitol (8 wks) vs. con	nbined calcipotriol + BMD	(4 wks); then calcipotriol (4 v	wks)		
Ortonne 2004	16/252	3/249	-	100.0 %	0.05 [0.02, 0.08
					-
Subtotal (95% CI)	252	249	•	100.0 %	0.05 [0.02, 0.08
	regimen), 3 (Complex regi	men)			
leterogeneity: not applicab					
Test for overall effect: $Z = 3$. ,				
est for subgroup difference	s: Chi ² = 15.88, df = 5 (P =	= 0.01), I ² =69%			
				L	
			-1 -0.5 0 0.5	1	

Analysis 13.9. Comparison 13 Vitamin D alone or in combination versus other treatments: complex regimens, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 13 Vitamin D alone or in combination versus other treatments: complex regimens

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D regimen	Complex regimen	Risk Difference	Weight	Risk Difference
	n/N	n/N	M- H,Random,95% Cl		-M H,Random,9 Cl
Calcipotriol (12 wks) vs.	combined calcipotriol + BN	1D (8 wks); then calcipotriol	(4 wks)		
Kragballe 2004	73/327	35/322		100.0 %	0.11 [0.06, 0.17]
Subtotal (95% CI)	327	322	*	100.0 %	0.11 [0.06, 0.17]
Total events: 73 (Vitamin D	regimen), 35 (Complex re	egimen)			
Heterogeneity: not applicat	ble				
Test for overall effect: $Z = 2$	3.97 (P = 0.000071)				
2 Calcipotriol (12 wks) vs.	combined calcipotriol + BM	1D (4 wks); then calcipotriol	(8 wks)		
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D i	regimen), 0 (Complex regir	men)			
Heterogeneity: not applicat	ble				
Test for overall effect: not a	pplicable				
3 Calcipotriol (12 wks) vs.	combined calcipotriol + BM	1D (4 wks); then calcipotriol	(w/dy) % combined calcipotric	ol + BMD (w/e) (8 wk	(s)
Kragballe 2004	73/327	37/322		100.0 %	0.11 [0.05, 0.17]
Subtotal (95% CI)	327	322	•	100.0 %	0.11 [0.05, 0.17]
Total events: 73 (Vitamin D	regimen), 37 (Complex re	egimen)			
Heterogeneity: not applicat	ble				
Test for overall effect: $Z = 2$	3.72 (P = 0.00020)				
4 Calcipotriol (6 wks) vs. cl	obetasol propionate (2 wks	s); then calcipotriol (4 wks)			
Austad 1998	4/49	3/49	-	100.0 %	0.02 [-0.08, 0.12]
Subtotal (95% CI)	49	49	•	100.0 %	0.02 [-0.08, 0.12]
Total events: 4 (Vitamin D i	8 , (1 8	men)			
Heterogeneity: not applicat					
Test for overall effect: $Z = 0$	· · · ·				
		· · · · ·	n calcipotriol twice daily (4 wk	,	
Wozel 2001	1/19	1/19		100.0 %	0.0 [-0.14, 0.14]
Subtotal (95% CI)	19	19	-	100.0 %	0.0 [-0.14, 0.14]
Total events: I (Vitamin D i	regimen), I (Complex regir	men)			
Heterogeneity: not applicat	ble				
		-(0.5 -0.25 0 0.25 0.	5	

(Continued ...)

Study or subgroup	Vitamin D regimen	Complex regimen	Risk Difference M-	Weight	Risk Difference M:
	n/N	n/N	H,Random,95% Cl		H,Random, C
Test for overall effect: Z = (0.0 (P = 1.0)				
6 Calcipotriol (6 wks) vs. ha	alometasone OM, calcipotrio	ol ON (2 wks); then calcipotri	iol twice daily (w/dy), halome	tasone (w/e) (2 wks);	then calcipotriol twice daily (
wks) Yang 2009	14/36	5/40		100.0 %	0.26 [0.07, 0.45
Subtotal (95% CI)	36	40	-	100.0 %	0.26 [0.07, 0.45]
Heterogeneity: not applicab Test for overall effect: Z = 2	2.73 (P = 0.0063)	men) wks); then calcipotriol twice o	daily (to wk 12) vs. calcitriol (DN, clobetasol propie	onate OM (2 to 4 wks); then
calcitriol twice daily (to w	rk 2)				
Lahfa 2003	3/64	4/61		100.0 %	-0.02 [-0.10, 0.06
Subtotal (95% CI)	64	61	+	100.0 %	-0.02 [-0.10, 0.06
Heterogeneity: not applicat Test for overall effect: Z = (0.45 (P = 0.65)	e ointment twice daily (8 wks)	vs. combined calcipotriol + I	BMD (4 wks); then ca	lcipotriol ointment twice dail
White 2006 (P)	32/373	43/379	-	100.0 %	-0.03 [-0.07, 0.02
Subtotal (95% CI)	373	379	•	100.0 %	-0.03 [-0.07, 0.02
Heterogeneity: not applicab Test for overall effect: Z = 9 Combined calcipotriol + calcipotriol + BMD (w/e) White 2006 (P)	I.27 (P = 0.20) BMD (4 wks); then placebo	o ointment twice daily (8 wks) 28/370) vs. combined calcipotriol +	BMD (4 wks); then ca	alcipotriol (w/dy) + combinec 0.01 [-0.03, 0.05
Subtotal (95% CI)	373	370	•	100.0 %	0.01 [-0.03, 0.05
Total events: 32 (Vitamin D Heterogeneity: not applicat Test for overall effect: Z = (regimen), 28 (Complex reg ole 0.51 (P = 0.61)	gimen)			
 Combined calcipotriol + 	. , , ,	otriol ointment twice daily (8	3 wks) vs. combined calcipotr	101 + BMD (4 wks); 1	then calcipotriol (w/dy) +
White 2006 (H)	43/379	28/370	-	100.0 %	0.04 [0.00, 0.08
Subtotal (95% CI)	379	370	•	100.0 %	0.04 [0.00, 0.08
	regimen), 28 (Complex reg ole			20010 /0	
I Combined calcipotriol +	+ BMD (8 wks); then calcipe	otriol (4 wks) vs. combined c	alcipotriol + BMD (4 wks); th	nen calcipotriol (w/dy) % combined calcipotriol +
BMD (w/e) (8 wks) Kragballe 2004	35/322	37/322	-	100.0 %	-0.01 [-0.05, 0.04
Subtotal (95% CI)	322	322	+	100.0 %	-0.01 [-0.05, 0.04
	regimen), 37 (Complex reg	gimen)			
Total events: 35 (Vitamin D		•			

Study or subgroup	Vitamin D regimen	Complex regimen	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
Heterogeneity: not applica	ble				
Test for overall effect: $Z =$	0.25 (P = 0.80)				
12 Tacalcitol (8 wks) vs. cc	ombined calcipotriol + BMD	(4 wks); then calcipotriol (4	wks)		
Ortonne 2004	32/252	16/249	· -	100.0 %	0.06 [0.01, 0.11]
Subtotal (95% CI)	252	249	•	100.0 %	0.06 [0.01, 0.11]
Total events: 32 (Vitamin E	D regimen), 16 (Complex re	gimen)			
Heterogeneity: not applica	ble				
Test for overall effect: Z =	2.40 (P = 0.016)				
Test for subgroup difference	ces: Chi ² = 35.10, df = 10 (F	P = 0.00), I ² =72%			
				1	
		-0	0.5 -0.25 0 0.25 0	0.5	

Favours vitamin D regimen Favours complex regimen

Analysis 14.1. Comparison 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks), Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks)

Outcome: I IAGI

Study or subgroup	Regimen I		Regimen 2		Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% CI
I Combined calcipotri	ol + BMD (52 wks)	vs. alternating: combine	ed calcipotriol + I	BMD (4 wks); then cal	cipotriol (4 wks)	
Kragballe 2006	104	1.74 (0.99)	104	1.83 (1.03)		-0.09 [-0.36, 0.18]
2 Combined calcipotri	ol + BMD (52 wks)	vs. combined calcipotr	iol + BMD (4 wk	s); then calcipotriol (48	3 wks)	
Kragballe 2006	104	1.74 (0.99)	89	1.92 (0.97)		-0.18 [-0.47, 0.10]
3 Alternating: combine	ed calcipotriol + BM	D (4 wks); then calcipo	triol (4 wks) vs. c	ombined calcipotriol -	- BMD (4 wks); then calcipotriol ((48 wks)
Kragballe 2006	104	1.83 (1.03)	89	1.92 (0.97)	- _	-0.09 [-0.37, 0.19]
					-1 -0.5 0 0.5 1	
				Fav	ours regimen I Favours regimer	n 2

Topical treatments for chronic plaque psoriasis (Review)

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Analysis 14.5. Comparison 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks), Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks)

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Regimen I		Regimen 2		Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% CI
I Combined calcipotri	iol + BMD (52 wks)	vs. alternating: combin	ed calcipotriol + I	BMD (4 wks); then a	calcipotriol (4 wks)	
Kragballe 2006	104	1.74 (0.99)	104	1.83 (1.03)	-	-0.09 [-0.36, 0.18]
2 Combined calcipotri	iol + BMD (52 wks)	vs. combined calcipotr	iol + BMD (4 wks	s); then calcipotriol ((48 wks)	
Kragballe 2006	104	1.74 (0.99)	89	1.92 (0.97)	-+	-0.18 [-0.47, 0.10]
3 Alternating: combine	ed calcipotriol + BM	⊃ (4 wks); then calcipo	otriol (4 wks) vs. c	ombined calcipotrio	I + BMD (4 wks); then calcipotriol	(48 wks)
Kragballe 2006	104	1.83 (1.03)	89	1.92 (0.97)	+	-0.09 [-0.37, 0.19]
					-4 -2 0 2 4	
					-7 -2 0 2 4	

Favours regimen 1 Favours regimen 2

Analysis 14.6. Comparison 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks), Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks)

Outcome: 6 Total withdrawals

Study or subgroup	Regimen I n/N	Regimen 2 n/N	Risk Difference M- H,Random,95% Cl	Risk Difference H,Random,95% Cl
				Ci
	, , ,		+ wks); then calcipotriol (4 wks)	
Kragballe 2006	64/212	56/213		0.04 [-0.05, 0.12]
2 Combined calcipotriol + E	3MD (52 wks) vs. combined ca	lcipotriol + BMD (4 wks); then	calcipotriol (48 wks)	
Kragballe 2006	64/212	70/209		-0.03 [-0.12, 0.06]
3 Alternating: combined cal	cipotriol + BMD (4 wks); then	calcipotriol (4 wks) vs. combine	ed calcipotriol + BMD (4 wks); then calcipo	otriol (48 wks)
Kragballe 2006	56/213	70/209		-0.07 [-0.16, 0.02]
			-0.5 -0.25 0 0.25 0.5	
			Favours regimen 1 Favours regimen 2	

Analysis 14.7. Comparison 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks), Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks)

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Regimen I	Regimen 2	Risk Difference M- H,Random,95%	Risk Difference M- H,Random,95%
	n/N	n/N	Cl	CI
Combined calcipotriol +	BMD (52 wks) vs. alternating: c	ombined calcipotriol + BMD (4	ł wks); then calcipotriol (4 wks)	
Kragballe 2006	14/212	11/213		0.01 [-0.03, 0.06]
2 Combined calcipotriol +	BMD (52 wks) vs. combined ca	Icipotriol + BMD (4 wks); then	calcipotriol (48 wks)	
Kragballe 2006	14/212	16/209		-0.01 [-0.06, 0.04]
3 Alternating: combined cal	cipotriol + BMD (4 wks); then	calcipotriol (4 wks) vs. combine	ed calcipotriol + BMD (4 wks); then calcip	potriol (48 wks)
Kragballe 2006	11/213	16/209		-0.02 [-0.07, 0.02]
			-0.2 -0.1 0 0.1 0.2	
			Favours regimen 1 Favours regimen 2	2

Analysis 14.8. Comparison 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks), Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks)

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Regimen I n/N	Regimen 2 n/N	Risk Difference H,Random,95%	Risk Difference H,Random,95%
	n/IN	n/IN	Cl	Cl
I Combined calcipotriol + I	BMD (52 wks) vs. alternating: c	ombined calcipotriol + BMD (4	wks); then calcipotriol (4 wks)	
Kragballe 2006	32/212	31/213	- <u>+</u> -	0.01 [-0.06, 0.07]
2 Combined calcipotriol + I	BMD (52 wks) vs. combined ca	lcipotriol + BMD (4 wks); then	calcipotriol (48 wks)	
Kragballe 2006	32/212	42/209		-0.05 [-0.12, 0.02]
3 Alternating: combined cal	cipotriol + BMD (4 wks); then	calcipotriol (4 wks) vs. combine	ed calcipotriol + BMD (4 wks); then calcip	potriol (48 wks)
Kragballe 2006	31/213	42/209		-0.06 [-0.13, 0.02]
			<u> </u>	
			-0.5 -0.25 0 0.25 0.5	
			Favours regimen 1 Favours regimen 2	2

Analysis 14.9. Comparison 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks), Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 14 Vitamin D alone or in combination versus other treatment: long-term studies (> 24 wks)

Outcome: 9 Adverse events (local)

Study or subgroup	Regimen I n/N	Regimen 2 n/N	Risk Difference M- H,Random,95% Cl	Risk Difference M- H,Random,95% Cl
Combined calcinotrial + B	MD (52 w/ks) vs. alternating: c	combined calcipotrial + BMD (4 wks); then calcipotriol (4 wks)	
Kragballe 2006	45/212	63/213		-0.08 [-0.17, 0.00]
2 Combined calcipotriol + B	MD (52 wks) vs. combined ca	alcipotriol + BMD (4 wks); ther	calcipotriol (48 wks)	
Kragballe 2006	45/212	78/209	· · · · · · · · · · · · · · · · · · ·	-0.16 [-0.25, -0.08]
3 Alternating: combined calc	ipotriol + BMD (4 wks); then	calcipotriol (4 wks) vs. combin	ed calcipotriol + BMD (4 wks); then ca	lcipotriol (48 wks)
Kragballe 2006	63/213	78/209		-0.08 [-0.17, 0.01]
			-0.5 -0.25 0 0.25 0.5	
			Favours regimen I Favours regime	n 2

Analysis 15.1. Comparison 15 Vitamin D analogues versus other treatment, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: I IAGI

St. Mea Weight Difference	Std. Mean Difference		ther treatment	С	Vitamin D analogue	Study or subgroup
IV,Random,95% (IV,Random,95% Cl	Mean(SD)	Ν	Mean(SD)	Ν	
						I Calcipotriol vs. coal tar
34.2 % 0.59 [0.04, 1.13	-	1.6 (1.01)	27	2.2 (1.01)	28	Alora-Palli 2010
32.0 % -1.01 [-1.77, -0.24	-	-1.8 (0.86)	15	-2.7 (0.88)	15	De Simone 1993
33.8 % -1.21 [-1.79, -0.62	-	-1.2 (0.96)	27	-2.3 (0.83)	27	Tham 1994
100.0 % -0.53 [-1.74, 0.68	•		69		70	Subtotal (95% CI)
			I ² =91%	2 (P = 0.00001)	= 0.86 (P = 0.39)	Heterogeneity: Tau ² = 1.0 Test for overall effect: Z = 2 Calcipotriol vs. coal tar
59.4 % -0.47 [-0.83, -0.11		-2.28 (0.92)	57	-2.66 (0.67)	, , ,	Pinheiro 1997
40.6 % -0.75 [-1.19, -0.32		-2.48 (1.24)	46	-3.41 (1.2)	41	Van de Kerkhof 2002a
100.0 % -0.59 [-0.87, -0.31	•		103		106	Subtotal (95% CI)
				$P = 0.33$; $I^2 = 0$	D; $Chi^2 = 0.94$, $df = 1$ (= 4.15 (P = 0.000034)	Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 3 Calcipotriol vs. nicotina
Not estimabl			0		0	Subtotal (95% CI)
				%, twice daily	applicable	Heterogeneity: not applic Test for overall effect: not 4 Calcipotriol vs. calcipoti
Not estimabl			0		0 able applicable teroid + salicylic acid	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: not 5 Calcipotriol vs. corticos
-0.06 [-0.33, 0.22		-2.65 (1.18)	100	-2.72 (1.27)	100	Scarpa 1994
100.0 % -0.06 [-0.33, 0.22	ł		100		= 0.40 (P = 0.69)	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z = 6 Calcipotriol vs. propylth
Not estimabl			0		0 able applicable	Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: not 7 Calcipotriol vs. tazarote
						Subtotal (95% CI)

(Continued . . .)

Study or subgroup	Vitamin D analogue	Oth	er treatment		Std. Mean Difference	Weight	(Continued) Std. Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	-	IV,Random,95% CI
Heterogeneity: not applie Test for overall effect: no 8 Calcipotriol vs. tacrolin	t applicable nus ointment						
Ortonne 2006	40	-3.25 (0.98)	84	-3.01 (1.12)	-	100.0 %	-0.22 [-0.60, 0.16]
Subtotal (95% CI) Heterogeneity: not applie Test for overall effect: Z 9 Calcipotriol vs. tazarote	cable = 1.15 (P = 0.25)	ne furnate cream	84		ł	100.0 %	-0.22 [-0.60, 0.16]
Subtotal (95% CI) Heterogeneity: not applie Test for overall effect: no	cable 0	le lui Gate cream	0				Not estimable
10 Calcipotriol vs. vitami Stuecker 2001		-2.08 (0.49)	13	-1.77 (0.6)	-	100.0 %	-0.55 [-1.33, 0.24]
Subtotal (95% CI) Heterogeneity: not applie Test for overall effect: Z	cable		13		•	1 00.0 %	-0.55 [-1.33, 0.24]
II Head-to-head vitamir	n D alone or in combina	ation: twice daily vs	OD				
Guenther 2002 (H)	234	-3.79 (0.97)	150	-3.59 (1)	•	51.6 %	-0.20 [-0.41, 0.00]
Kragballe 1998b	172	-2.98 (1.23)	172	-2.63 (1.34)	•	48.4 %	-0.27 [-0.48, -0.06]
Subtotal (95% CI) Heterogeneity: $Tau^2 = 0$. Test for overall effect: Z	.0; $Chi^2 = 0.20$, $df = 1$	(P = 0.65); I ² =0.0%	322		•	100.0 %	-0.24 [-0.38, -0.09]
12 Head-to-head vitamir Subtotal (95% CI) Heterogeneity: not applie Test for overall effect: no	0 0 cable	ation: no occlusion [.]	vs. occlusion 0				Not estimable
				-10 Favours vitamin D	-5 0 5 analogue Favours ot	10 her treatment	

Analysis 15.2. Comparison 15 Vitamin D analogues versus other treatment, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: 2 TSS

Study or subgroup	Vitamin D analogue	Othe	er treatment		Std. Mean Difference	Weight	Std Mear Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% C
I Calcipotriol vs. coal tar							
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applica	ble						
Test for overall effect: not a	applicable						
2 Calcipotriol vs. coal tar p	oolytherapy						
Pinheiro 1997	69	2.7 (2.14)	63	3.8 (2.14)	-	100.0 %	-0.51 [-0.86, -0.16
Subtotal (95% CI)	69		63		•	100.0 %	-0.51 [-0.86, -0.16
Heterogeneity: not applica	ble						
Test for overall effect: Z =	2.88 (P = 0.0039)						
3 Calcipotriol vs. nicotinam	nide 1.4%, twice daily						
Levine 2010 (H)	48	3.52 (1.95)	48	3.69 (1.93)	-	100.0 %	-0.09 [-0.49, 0.31
Subtotal (95% CI)	48		48		•	100.0 %	-0.09 [-0.49, 0.31
Heterogeneity: not applica	ble						
Test for overall effect: Z =	0.43 (P = 0.67)						
4 Calcipotriol vs. calcipotri	ol+nicotinamide (0.0	5%, 0.1%, 0.7%, or	1.4%), twice c	laily			
Levine 2010 (H)	48	3.52 (1.95)	144	3.15 (1.94)	•	100.0 %	0.19 [-0.14, 0.52
Subtotal (95% CI)	48		144		•	100.0 %	0.19 [-0.14, 0.52
Heterogeneity: not applica						,	
Test for overall effect: Z =							
5 Calcipotriol vs. corticost	. ,						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applica	ble						
Test for overall effect: not a	applicable						
6 Calcipotriol vs. propylthi	ouracil cream						
Subtotal (95% CI)	0		0				Not estimabl
Heterogeneity: not applica	ble						
Test for overall effect: not a	applicable						
7 Calcipotriol vs. tacrolimu	is ointment						
Ortonne 2006	40	3.25 (0.98)	84	3.01 (1.12)	-	52.1 %	0.22 [-0.16, 0.60
Zonneveld 1998 (H)	23	2.6 (2.14)	24	4.7 (2.14)	-	47.9 %	-0.96 [-1.57, -0.36
Subtotal (95% CI)	63		108		•	100.0 %	-0.35 [-1.51, 0.81
Heterogeneity: $Tau^2 = 0.64$	4; Chi ² = 10.58, df =	$ (P = 0.00); ^2 =$	91%				
Test for overall effect: Z =	0.59 (P = 0.56)	- *					
				-10	-5 0 5	10	
				Favours vitamin D	analogue Favours of	her treatment	
					-		(Continued

(Continued \dots)

					Std.		(Continued Std.
Study or subgroup Vitamin D	analogue	Othe	r treatment		Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	IV,Random,95% CI	, reight	IV,Random,95% CI
8 Calcipotriol vs. tazarotene							
Han 2001	101	3.91 (6.08)	98	4.18 (4.39)	•	100.0 %	-0.05 [-0.33, 0.23]
Subtotal (95% CI)	101		98		•	100.0 %	-0.05 [-0.33, 0.23]
Heterogeneity: not applicable							
Test for overall effect: $Z = 0.36$ (P =	0.72)						
9 Calcipotriol vs. tazarotene gel plus	mometasc	ne furoate cream					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable							
Test for overall effect: not applicable							
10 Calcipotriol vs. vitamin B12 crean	า						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable							
Test for overall effect: not applicable							
II Head-to-head vitamin D alone or	r in combin	ation: twice daily vs	OD				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable							
Test for overall effect: not applicable							
12 Head-to-head vitamin D alone or	in combin	ation: no occlusion v	/s. occlusion				
Bourke 1993b	19	-3.1 (2.6)	19	-5.2 (2.6)	-	48.7 %	0.79 [0.13, 1.45]
Hinds n 2006 (H)	4	5.2 (3.3)	95	9.7 (4.8)		51.3 %	- . [- .40, -0.8]
Subtotal (95% CI)	133		114		+	100.0 %	-0.18 [-2.04, 1.68]
Heterogeneity: Tau ² = 1.73 ; Chi ² =	26.34, df =	(P<0.0000); ² =	=96%				
Test for overall effect: $Z = 0.19$ (P =	0.85)						
				-10	-5 0 5	10	
				Favours vitamin D	analogue Favours	other treatment	

Analysis 15.3. Comparison 15 Vitamin D analogues versus other treatment, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: 3 PASI

Sta Mea Differenc IV,Random,95% (Weight	Std. Mean Difference IV.Random,95% Cl	Mean(SD)	Other treatment	Mean(SD)	Vitamin D analogue N	Study or subgroup
							I Calcipotriol vs. coal tar
0.64 [0.10, 1.18	50.1 %		-0.58 (0.37)	27	-0.37 (0.27)	28	Alora-Palli 2010
-0.84 [-1.39, -0.28	49.9 %		4.5 (3.6)	27	2 (2.1)	27	Tham 1994
-0.10 [-1.54, 1.35	100.0 %		-	54		55	Subtotal (95% CI)
				0); I ² =93%	I (P = 0.0002	= 0.13 (P = 0.90)	Heterogeneity: Tau ² = 1. Test for overall effect: Z = 2 Calcipotriol vs. coal tar
-0.63 [-1.06, -0.20	100.0 %		-0.36 (0.36)	46	-0.57 (0.29)	a 41	Van de Kerkhof 2002a
-0.63 [-1.06, -0.20]	100.0 %	•		46		41	Subtotal (95% CI)
						= 2.87 (P = 0.0041)	Heterogeneity: not applic Test for overall effect: Z = 3 Calcipotriol vs. nicotina
Not estimable				0		0	Subtotal (95% CI)
						able	Heterogeneity: not applic
						t applicable	Test for overall effect: not
					%, twice daily		4 Calcipotriol vs. calcipot
Not estimable				0			Subtotal (95% CI)
							Heterogeneity: not applic
							Test for overall effect: not
-0.05 [-0.36, 0.26	100.0 %	-	2.78 (3.61)	80	2.6 (3.61)	80	5 Calcipotriol vs. corticos Crosti 1997
-0.05 [-0.36, 0.26	100.0 %	•	~ /	80	× /	80	Subtotal (95% CI)
10.09 [10.90, 0.20	100.0 /0					able = 0.31 (P = 0.75)	Heterogeneity: not applic Test for overall effect: Z = 6 Calcipotriol vs. propylth
-2.24 [-3.23, -1.25	100.0 %	_	3.83 (1.31)	13	1.11 (1.04)	4	Sanchez 2001
2.24 [-3.23, -1.25	100.0 %	-		13		14	Subtotal (95% CI) Heterogeneity: not applic
							Test for overall effect: Z =
						us ointment	7 Calcipotriol vs. tacrolim
Not estimable				0		0	Subtotal (95% CI)
							Heterogeneity: not applic Test for overall effect: not

(Continued . . .)

(Continue Sti Mea Differenc IV,Random,95% (Weight	Std. Mean Difference dom,95% Cl		Mean(SD)	treatment N	Othe Mean(SD)	n D analogue N	Study or subgroup Vitamin
				. /		. ,		8 Calcipotriol vs. tazarotene
Not estimabl					0		0	Subtotal (95% CI)
								Heterogeneity: not applicable
							ble	Test for overall effect: not applicable
						e furoate cream		9 Calcipotriol vs. tazarotene gel plu
Not estimable					0		0	Subtotal (95% CI)
								Heterogeneity: not applicable
							ble	Test for overall effect: not applicable
							am	10 Calcipotriol vs. vitamin B12 crea
-0.01 [-0.78, 0.75	100.0 %	-		0.81 (0.69)	13	0.8 (0.66)	13	Stuecker 2001
-0.01 [-0.78, 0.75	100.0 %	-			13		13	Subtotal (95% CI)
								Heterogeneity: not applicable
							= 0.97)	Test for overall effect: Z = 0.04 (P =
					D	tion: twice daily vs	or in combina	II Head-to-head vitamin D alone o
-0.10 [-0.34, 0.15	26.8 %	•	-	1.1 (1.4)	130	0.97 (1.3)	130	Baiocchi 1997
-0.12 [-0.32, 0.09	37.7 %	-	-	3 (2.5)	150	2.7 (2.5)	234	Guenther 2002 (H)
-0.15 [-0.36, 0.06	35.5 %	-	-	4.58 (3.93)	173	4.04 (3.39)	172	Kragballe 1998b
-0.12 [-0.25, 0.00	100.0 %	•	•		453		536	Subtotal (95% CI)
					-50	$P = 0.95$); $ ^2 = 0.0\%$		Heterogeneity: $Tau^2 = 0.0$; $Chi^2 = 0$
						,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		Test for overall effect: Z = 1.91 (P =
								12 Head-to-head vitamin D alone o
Not estimabl					000000000000000000000000000000000000000	lion. no occiusion v	0 11 0011011a	Subtotal (95% CI)
					Ŭ		v	Heterogeneity: not applicable
							ble	Test for overall effect: not applicable

Favours vitamin D analogue Favour

Favours other treatment

Analysis 15.4. Comparison 15 Vitamin D analogues versus other treatment, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: 4 PAGI

St: Mea Neight Differenc IV,Random,95% (Std. Mean rence n,95% Cl	Diffe IV,Rando	Mean(SD)	Other treatment N	Mean(SD)	Vitamin D analogue N	Study or subgroup
							I Calcipotriol vs. coal tar
00.0 % -1.51 [-2.12, -0.90			-1.11 (0.85) +	27	-2.44 (0.89)	27	Tham 1994
0.0 % -1.51 [-2.12, -0.90	10	-	-	27		27	Subtotal (95% CI)
						ble	Heterogeneity: not applic
						4.84 (P < 0.00001)	Test for overall effect: Z =
						oolytherapy	2 Calcipotriol vs. coal tar
-0.56 [-0.99, -0.13			-2.57 (1.36)	46	-3.29 (1.19)	41	Van de Kerkhof 2002a
0.0 % -0.56 [-0.99, -0.13	10	-		46		41	Subtotal (95% CI)
						ble	Heterogeneity: not applic
						2.54 (P = 0.011)	Test for overall effect: Z =
						nide 1.4%, twice daily	3 Calcipotriol vs. nicotinal
Not estimabl				0		0	Subtotal (95% CI)
						ble	Heterogeneity: not applic
						applicable	Test for overall effect: not
					%, twice daily		4 Calcipotriol vs. calcipotr
Not estimabl				0		0	Subtotal (95% CI)
							Heterogeneity: not applic
							Test for overall effect: not
						,	5 Calcipotriol vs. corticos
00.0 % -0.49 [-0.79, -0.20			-2.87 (0.94)	97	-3.29 (0.73)	89	Scarpa 1994
0.0 % -0.49 [-0.79, -0.20	10	•		97		89	Subtotal (95% CI)
						ble	Heterogeneity: not applic
						3.32 (P = 0.00091)	Test for overall effect: Z =
							6 Calcipotriol vs. propylth
Not estimabl				0		0	Subtotal (95% CI)
							Heterogeneity: not applic
							Test for overall effect: not
			200 (112)	0.4	2.22 (1.05)		7 Calcipotriol vs. tacrolim
-0.13 [-0.51, 0.24			-3.08 (1.13)	84	-3.23 (1.05)	40	Ortonne 2006
0.0 % -0.13 [-0.51, 0.24	10	-		84		40	Subtotal (95% CI)
							Heterogeneity: not applic
						. ,	Test for overall effect: Z =
						ie	8 Calcipotriol vs. tazarote
	<u> </u>						

(Continued \dots)

Study or subgroup	Vitamin D analogue		r treatment		Std. Mean Difference	Weight	(Continued) Std. Mean Difference
T 2005	N 10	Mean(SD)	N	Mean(SD)	IV,Random,95% CI	100.0.00	IV,Random,95% C
Tzung 2005	19	-3.16 (0.69)	19	-2.89 (0.81)	-	100.0 %	-0.35 [-0.99, 0.29]
Subtotal (95% CI)	19		19		-	100.0 %	-0.35 [-0.99, 0.29]
Heterogeneity: not applical							
Test for overall effect: $Z =$,						
9 Calcipotriol vs. tazaroten	0 1	ne furoate cream					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applical							
Test for overall effect: not a	applicable						
10 Calcipotriol vs. vitamin l	B12 cream						
Stuecker 2001	13	-2.08 (0.49)	13	-1.77 (0.6)		100.0 %	-0.55 [-1.33, 0.24]
Subtotal (95% CI)	13		13			100.0 %	-0.55 [-1.33, 0.24]
Heterogeneity: not applical	ole						
Test for overall effect: Z =	1.37 (P = 0.17)						
II Head-to-head vitamin E) alone or in combin	ation: twice daily vs (DD				
Subtotal (95% CI)	0	,	0				Not estimable
Heterogeneity: not applical	ole						
Test for overall effect: not a	applicable						
12 Head-to-head vitamin [) alone or in combin	ation: no occlusion v	s. occlusion				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applical	ole						
Test for overall effect: not a	applicable						
						_1	
				-2	-1 0 1	2	
				Favours vitamin D	analogue Favours of	her treatment	

Analysis 15.5. Comparison 15 Vitamin D analogues versus other treatment, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D analogue N	Mean(SD)	Other treatment	Mean(SD)	Std. Mean Difference IV.Random,95% Cl	Weight	Std. Mean Difference IV.Random,95% CI
		T lean(SD)	11	Tiedii(5D)	14,1 & 10011,7 576 CI		17,1 and 011,7378 CI
l Calcipotriol vs. coal tar Alora-Palli 2010	28	2.2 (1.01)	27	1.6 (1.01)		34.2 %	0.59 [0.04, 1.13]
De Simone 1993	15	-2.7 (0.88)	15	-1.8 (0.86)		32.0 %	-1.01 [-1.77, -0.24]
		. ,					
Tham 1994	27	-2.3 (0.83)	27	-1.2 (0.96)		33.8 %	-1.21 [-1.79, -0.62]
Subtotal (95% CI)			69		-	100.0 %	-0.53 [-1.74, 0.68]
Heterogeneity: $Tau^2 = 1$.		2 (P = 0.0000	$); ^2 = 9 \%$				
Test for overall effect: Z = 2 Calcipotriol vs. coal tar	· · · ·						
Pinheiro 1997	. , .,	-2.66 (0.67)	57	-2.28 (0.92)	-	59.4 %	-0.47 [-0.83, -0.11]
Van de Kerkhof 2002		-3.41 (1.2)		-2.48 (1.24)		40.6 %	-0.75 [-1.19, -0.32]
		-3.41 (1.2)		-2.40 (1.24)	_		
Subtotal (95% CI)		(0.000) 12	103		•	100.0 %	-0.59 [-0.87, -0.31]
Heterogeneity: $Tau^2 = 0$.		· /	=0.0%				
Test for overall effect: Z = 3 Calcipotriol vs. nicotina	· · · · · · · · · · · · · · · · · · ·						
Levine 2010 (H)	48	3.52 (1.95)	48	3.69 (1.93)		100.0 %	-0.09 [-0.49, 0.31]
Subtotal (95% CI)	48		48		•		-0.09 [-0.49, 0.31]
Heterogeneity: not applie			40			100.0 /0	-0.07 [-0.17, 0.51]
Test for overall effect: Z							
4 Calcipotriol vs. calcipot	· · · · ·	05%, 0.1%, 0.7	%, or 1.4%), twice	daily			
Levine 2010 (H)	48	3.52 (1.95)	44	3.15 (1.94)	••	100.0 %	0.19 [-0.14, 0.52]
Subtotal (95% CI)	48		144		•	100.0 %	0.19 [-0.14, 0.52]
Heterogeneity: not applie	cable						
Test for overall effect: Z	= 1.14 (P = 0.26)						
5 Calcipotriol vs. cortico:	steroid + salicylic acid						
Crosti 1997	80	2.6 (3.61)	80	2.78 (3.61)	=	44.4 %	-0.05 [-0.36, 0.26]
Scarpa 1994	100	-2.72 (1.27)	100	-2.65 (1.18)	-	55.6 %	-0.06 [-0.33, 0.22]
Subtotal (95% CI)	180		180		+	100.0 %	-0.05 [-0.26, 0.15]
Heterogeneity: $Tau^2 = 0$.	.0; $Chi^2 = 0.00$, $df = 1$	(P = 0.97); I ²	=0.0%				
Test for overall effect: Z	= 0.51 (P = 0.61)						
6 Calcipotriol vs. propyltl	hiouracil cream						
						1	
				_4	ł -2 0 2	4	
				Favours vitamin	D analogue Favours ot	her treatment	,

(Continued . . .)

Study or subgroup	Vitamin D analogue	Oth	ner treatment		Std. Mean Difference	Weight	(Continued Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% C
Sanchez 2001	14	1.11 (1.04)	13	3.83 (1.31)		100.0 %	-2.24 [-3.23, -1.25]
Subtotal (95% CI)	14		13		-	100.0 %	-2.24 [-3.23, -1.25]
Heterogeneity: not applic Test for overall effect: Z :							
7 Calcipotriol vs. tacrolim	· · · · · · · · · · · · · · · · · · ·						
Ortonne 2006	40	-3.25 (0.98)	84	-3.01 (1.12)	-	55.3 %	-0.22 [-0.60, 0.16
Zonneveld 1998 (H)	23	2.6 (2.14)	24	4.7 (2.14)		44.7 %	-0.96 [-1.57, -0.36]
Subtotal (95% CI)	63		108		-	100.0 %	-0.55 [-1.28, 0.17]
Heterogeneity: Tau ² = 0. Test for overall effect: Z = 8 Calcipotriol vs. tazarote	= 1.50 (P = 0.13) ene					040.07	
Han 2001	101	3.91 (6.08)	98	4.18 (4.39)		84.2 %	-0.05 [-0.33, 0.23]
Tzung 2005	19	-3.16 (0.69)	19	-2.89 (0.81)		15.8 %	-0.35 [-0.99, 0.29]
Subtotal (95% CI) Heterogeneity: Tau ² = 0. Test for overall effect: Z = 9 Calcipotriol vs. tazarote	0; Chi ² = 0.71, df = 1 (= 0.75 (P = 0.45)		117 %		•	100.0 %	-0.10 [-0.35, 0.16]
Subtotal (95% CI) Heterogeneity: not applie Test for overall effect: no	able		0				Not estimable
10 Calcipotriol vs. vitamir					_		
Stuecker 2001	13	-2.08 (0.49)	13	-1.77 (0.6)		100.0 %	-0.55 [-1.33, 0.24]
Subtotal (95% CI) Heterogeneity: not applie Test for overall effect: Z =	able		13		-	100.0 %	-0.55 [-1.33, 0.24]
II Head-to-head vitamin	D alone or in combina	ation: twice daily v	s OD				
Baiocchi 1997	130	0.97 (1.3)	130	. (.4)	•	26.9 %	-0.10 [-0.34, 0.15]
Guenther 2002 (H)	234	-3.79 (0.97)	150	-3.59 (1)	•	37.7 %	-0.20 [-0.41, 0.00]
Kragballe 1998b	172	-2.98 (1.23)	172	-2.63 (1.34)	=	35.3 %	-0.27 [-0.48, -0.06]
Subtotal (95% CI) Heterogeneity: Tau ² = 0. Test for overall effect: Z =	0; $Chi^2 = 1.14$, $df = 2$ ($(P = 0.57); I^2 = 0.0)$	452		•	100.0 %	-0.20 [-0.32, -0.07]
12 Head-to-head vitamin				52.00		407.0/	
Bourke 1993b	19	-3.1 (2.6)	19	-5.2 (2.6)	_ -	48.7 %	0.79 [0.13, 1.45]
Hinds n 2006 (H)	114	5.2 (3.3)	95	9.7 (4.8)	-	51.3 %	-1.11 [-1.40, -0.81]
Subtotal (95% CI) Heterogeneity: Tau ² = 1. Test for overall effect: Z =		I (P<0.00001); I ²	114 =96%			100.0 %	-0.18 [-2.04, 1.68]

(Continued . . .)

Study or subgroup	Vitamin D analogue	Other tre	eatment		Diff	Std. Mean erence	Weight	(Continued) Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Rando	om,95% Cl		IV,Random,95% CI
Test for subgroup differe	nces: Chi ² = 33.90, df =	10 (P = 0.00), I ² =719	%					
				1	1		ı	
				-4	-2 0) 2 4	4	
				Favours vitamin D a	analogue	Favours othe	er treatment	

Analysis 15.6. Comparison 15 Vitamin D analogues versus other treatment, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,99 Cl
l Calcipotriol vs. coal tar					
Alora-Palli 2010	2/30	3/30		54.2 %	-0.03 [-0.17, 0.11]
Tham 1994	3/30	3/30		45.8 %	0.0 [-0.15, 0.15]
Subtotal (95% CI)	60	60		100.0 %	-0.02 [-0.12, 0.08]
	analogue), 6 (Other treatment Chi ² = 0.10, df = 1 (P = 0.7	,			
2 Calcipotriol vs. coal tar po					
Pinheiro 1997	4/69	6/63		60.5 %	-0.04 [-0.13, 0.05]
Van de Kerkhof 2002a	3/41	4/47	_	39.5 %	-0.01 [-0.12, 0.10]
Subtotal (95% CI)	110	110	-	100.0 %	-0.03 [-0.10, 0.04]
Total events: 7 (Vitamin D a	analogue), 10 (Other treatm	ent)			
Heterogeneity: $Tau^2 = 0.0;$	$Chi^2 = 0.12, df = 1 (P = 0.7)$	3); I ² =0.0%			
Test for overall effect: $Z = 0$	0.75 (P = 0.45)				
3 Calcipotriol vs. nicotinami	ide 1.4%, twice daily				
Levine 2010 (H)	2/48	1/48		100.0 %	0.02 [-0.05, 0.09]
Subtotal (95% CI)	48	48		100.0 %	0.02 [-0.05, 0.09]
		-	0.2 -0.1 0 0.1 0.2	2	
		Favours vitam	in D analogue Favours other	treatment	

(Continued . . .)

Topical treatments for chronic plaque psoriasis (Review)

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Difference Difference Study or subgroup Vitamin D analogue Other treatment Weight M-M-H,Random,95% H,Random,95% n/N n/N ĆΓ Ć Total events: 2 (Vitamin D analogue), I (Other treatment) Heterogeneity: not applicable Test for overall effect: Z = 0.59 (P = 0.56) 4 Calcipotriol vs. calcipotriol + nicotinamide (0.05%, 0.1%, 0.7%, or 1.4%), twice daily Levine 2010 (H) 2/48 5/144 100.0 % 0.01 [-0.06, 0.07] Subtotal (95% CI) 48 144 100.0 % 0.01 [-0.06, 0.07] Total events: 2 (Vitamin D analogue), 5 (Other treatment) Heterogeneity: not applicable Test for overall effect: Z = 0.21 (P = 0.83) 5 Calcipotriol vs. corticosteroid + salicylic acid Crosti 1997 20/80 17/80 100.0 % 0.04 [-0.09, 0.17] Subtotal (95% CI) 80 80 100.0 % 0.04 [-0.09, 0.17] Total events: 20 (Vitamin D analogue), 17 (Other treatment) Heterogeneity: not applicable Test for overall effect: Z = 0.56 (P = 0.57) 6 Calcipotriol vs. propylthiouracil cream Sanchez 2001 2/14 1/14 ----100.0 % 0.07 [-0.16, 0.30] Subtotal (95% CI) 14 14 100.0 % 0.07 [-0.16, 0.30] Total events: 2 (Vitamin D analogue), I (Other treatment) Heterogeneity: not applicable Test for overall effect: Z = 0.62 (P = 0.54) 7 Calcipotriol vs. tacrolimus ointment Ortonne 2006 3/40 17/84 100.0 % -0.13 [-0.25, -0.01] Subtotal (95% CI) -0.13 [-0.25, -0.01] 40 84 100.0 % Total events: 3 (Vitamin D analogue), 17 (Other treatment) Heterogeneity: not applicable Test for overall effect: Z = 2.11 (P = 0.035) 8 Calcipotriol vs. tazarotene Han 2001 2/103 7/105 94.1 % -0.05 [-0.10, 0.01] Tzung 2005 4/23 0.0 [-0.22, 0.22] 4/23 5.9 % Subtotal (95% CI) 126 128 100.0 % -0.04 [-0.10, 0.01] Total events: 6 (Vitamin D analogue), 11 (Other treatment) Heterogeneity: Tau² = 0.0; Chi² = 0.21, df = 1 (P = 0.64); l² = 0.0% Test for overall effect: Z = 1.64 (P = 0.10) 9 Calcipotriol vs. tazarotene gel plus mometasone furoate cream Guenther 2000 100.0 % -0.03 [-0.15, 0.08] 6/60 8/60 Subtotal (95% CI) 60 60 100.0 % -0.03 [-0.15, 0.08] Total events: 6 (Vitamin D analogue), 8 (Other treatment) Heterogeneity: not applicable Test for overall effect: Z = 0.57 (P = 0.57)

-0.2 -0.1

Favours vitamin D analogue

0 0.1 0.2

Favours other treatment

Risk

(Continued . . .)

(... Continued)

Risk

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M- H,Random,95%	Weight	(Continued) Risk Difference M- H,Random,95%
	n/N	n/N	Cl		CI
10 Calcipotriol vs. vitamin B					
Stuecker 2001	0/13	0/13		100.0 %	0.0 [-0.14, 0.14]
Subtotal (95% CI)	13	13		100.0 %	0.0 [-0.14, 0.14]
	analogue), 0 (Other treatmen	nt)			
Heterogeneity: not applicab					
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
	alone or in combination: tw	,			
Baiocchi 1997	33/132	34/132		13.3 %	-0.01 [-0.11, 0.10]
Guenther 2002 (H)	19/237	11/152		50.8 %	0.01 [-0.05, 0.06]
Kragballe 1998b	19/174	17/174	_ _	35.9 %	0.01 [-0.05, 0.08]
Subtotal (95% CI)	543	458	-	100.0 %	0.01 [-0.03, 0.05]
	analogue), 62 (Other treatm Chi ² = 0.11, df = 2 (P = 0.9				
Test for overall effect: $Z = 0$					
12 Head-to-head vitamin D	alone or in combination: no	occlusion vs. occlusion			
Subtotal (95% CI)	0	0			Not estimable
	analogue), 0 (Other treatme	nt)			
Heterogeneity: not applicab					
Test for overall effect: not a	pplicable				
		-0	.2 -0.1 0 0.1 0.2)	
		Favours vitamir			

Analysis 15.7. Comparison 15 Vitamin D analogues versus other treatment, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

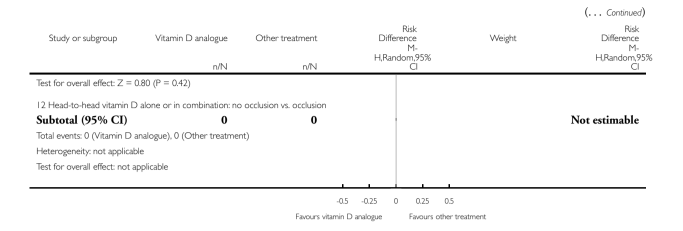
Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
l Calcipotriol vs. coal tar					
Alora-Palli 2010	0/30	0/30	#	66.1 %	0.0 [-0.06, 0.06]
Tham 1994	1/30	0/30		33.9 %	0.03 [-0.05, 0.12]
Subtotal (95% CI)	60	60	+	100.0 %	0.01 [-0.04, 0.06]
,	,	·			
Pinheiro 1997	1/65	3/57	-	64.3 %	-0.04 [-0.10, 0.03]
Van de Kerkhof 2002a	2/41	2/47		35.7 %	0.01 [-0.08, 0.09]
Subtotal (95% CI)	106	104	•	100.0 %	-0.02 [-0.07, 0.03]
Test for overall effect: Z = 0 3 Calcipotriol vs. nicotinamic	de 1.4%, twice daily	, 	_	100.0 %	0.0 [-0.04 0.04
Levine 2010 (H)	0/48	0/48	-	100.0 %	0.0 [-0.04, 0.04]
Subtotal (95% CI)	48	48	+	100.0 %	0.0 [-0.04, 0.04]
Heterogeneity: not applicabl Test for overall effect: Z = 0			у		
Levine 2010 (H)	0/48	0/144	-	100.0 %	0.0 [-0.03, 0.03]
Subtotal (95% CI) Total events: 0 (Vitamin D ar Heterogeneity: not applicabl Test for overall effect: Z = 0 5 Calcipotriol vs. corticoster	0.0 (P = 1.0)	144 t)	•	100.0 %	0.0 [-0.03, 0.03]
Crosti 1997	4/80	0/80	-	100.0 %	0.05 [0.00, 0.10]
Subtotal (95% CI)	80	80	•	100.0 %	0.05 [0.00, 0.10]
· · · · ·	nalogue), 0 (Other treatmen				
		-0 Favours vitamin			(Continued

					(Continued
Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
test for overall effect: $Z = 1.8$	36 (P = 0.063)				
Calcipotriol vs. propylthiour	racil cream				
Sanchez 2001	2/14	/ 4		100.0 %	0.07 [-0.16, 0.30]
Subtotal (95% CI)	14	14		100.0 %	0.07 [-0.16, 0.30]
otal events: 2 (Vitamin D an	alogue), I (Other treatmer	nt)			
Heterogeneity: not applicable					
est for overall effect: $Z = 0.6$	62 (P = 0.54)				
' Calcipotriol vs. tacrolimus c					
Ortonne 2006	3/40	5/84		100.0 %	0.02 [-0.08, 0.11]
Subtotal (95% CI)	40	84	-	100.0 %	0.02 [-0.08, 0.11]
otal events: 3 (Vitamin D an		nt)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.3$	32 (P = 0.75)				
Calcipotriol vs. tazarotene	2/102	E LLOE	_	70.0.9/	
Han 2001	3/103	5/105	-	70.0 %	-0.02 [-0.07, 0.03]
Tzung 2005	0/23	3/23		30.0 %	-0.13 [-0.28, 0.02]
Subtotal (95% CI)	126	128	-	100.0 %	-0.05 [-0.16, 0.05]
 Fest for overall effect: Z = 0.9 Calcipotriol vs. tazarotene g Guenther 2000 		te cream 4/60	-	100.0 %	-0.02 [-0.10, 0.07]
Subtatel (05% CI)	60	60	-	100 0 04	
Subtotal (95% CI) otal events: 3 (Vitamin D an				100.0 %	-0.02 [-0.10, 0.07]
leterogeneity: not applicable est for overall effect: $Z = 0.3$		ιτ <i>)</i>			
0 Calcipotriol vs. vitamin BI		0/12			
Stuecker 2001	0/13	0/13		100.0 %	0.0 [-0.14, 0.14]
Subtotal (95% CI) otal events: 0 (Vitamin D an Heterogeneity: not applicable oest for overall effect: Z = 0.0		13		100.0 %	0.0 [-0.14, 0.14]
I Head-to-head vitamin D a	lone or in combination: tw	ice daily vs OD			
Baiocchi 1997	7/132	6/132	-	5.8 %	0.01 [-0.04, 0.06]
Guenther 2002 (H)	1/235	0/151	•	85.0 %	0.00 [-0.01, 0.02]
Kragballe 1998b	8/174	6/174	+	9.3 %	0.01 [-0.03, 0.05]
	541	457	•	100.0 %	0.01 [-0.01, 0.02]

Favours vitamin D analogue Favours other treatment

(Continued . . .)



Analysis 15.8. Comparison 15 Vitamin D analogues versus other treatment, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
l Calcipotriol vs. coal tar					
Tham 1994	0/30	0/30	-	100.0 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	30	30	•	100.0 %	0.0 [-0.06, 0.06]
Total events: 0 (Vitamin D a	analogue), 0 (Other treatmer	nt)			
Heterogeneity: not applicat	ble				
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
2 Calcipotriol vs. coal tar p	olytherapy				
Van de Kerkhof 2002a	1/41	1/47	-	100.0 %	0.00 [-0.06, 0.07]
Subtotal (95% CI)	41	47	+	100.0 %	0.00 [-0.06, 0.07]
Total events: I (Vitamin D a	analogue), I (Other treatmer	nt)			
Heterogeneity: not applicat	ble				
Test for overall effect: $Z = 0$	0.10 (P = 0.92)				
			-I -0.5 0 0.5 I		
		Favours vitar	nin D analogue Favours other	treatment	
					(Continued)

Topical treatments for chronic plaque psoriasis (Review)

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(... Continued)

					(Continue
Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M-	Weight	Risl Difference M
	n/N	n/N	H,Random,95% Cl		H,Random, C
3 Calcipotriol vs. nicotinamic					
Levine 2010 (H)	0/48	0/48	-	100.0 %	0.0 [-0.04, 0.04
Subtotal (95% CI)	48	48	•	100.0 %	0.0 [-0.04, 0.04
Total events: 0 (Vitamin D ar					
Heterogeneity: not applicable	. , , ,	,			
Test for overall effect: $Z = 0$.	0 (P = 1.0)				
4 Calcipotriol vs. calcipotriol	+ nicotinamide (0.05%, 0.1	%, 0.7%, or 1.4%), twice dail	у		
Levine 2010 (H)	0/48	0/144		100.0 %	0.0 [-0.03, 0.03
Subtotal (95% CI)	48	144	+	100.0 %	0.0 [-0.03, 0.03
Total events: 0 (Vitamin D ar	nalogue), 0 (Other treatme	nt)			
Heterogeneity: not applicable	. , , ,	,			
Test for overall effect: $Z = 0$.					
5 Calcipotriol vs. corticosten	. ,				
Crosti 1997	, 1/80	3/80		100.0 %	-0.02 [-0.07, 0.02
Subtotal (95% CI)	80	80	•	100.0 %	-0.02 [-0.07, 0.02]
Total events: I (Vitamin D ar				10000 /0	0.02[0.07, 0.02
Heterogeneity: not applicable					
Test for overall effect: $Z = 1$.					
6 Calcipotriol vs. propylthiou	. ,				
Sanchez 2001	0/14	1/14		100.0 %	-0.07 [-0.25, 0.11
Subtotal (95% CI)	14	14	•	100.0 %	-0.07 [-0.25, 0.11
Total events: 0 (Vitamin D ar				100.0 /0	-0.07 [-0.29, 0.11
Heterogeneity: not applicable	. , , ,				
Test for overall effect: $Z = 0$.					
7 Calcipotriol vs. tacrolimus	. ,				
Ortonne 2006	0/40	2/84	+	100.0 %	-0.02 [-0.07, 0.03
			Ţ		-
Subtotal (95% CI)	40	84	•	100.0 %	-0.02 [-0.07, 0.03
Total events: 0 (Vitamin D ar	8 / (nt)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0$.	75 (P = 0.34)				
8 Calcipotriol vs. tazarotene Han 2001	0/103	0/105		100.0 %	
			T		0.0 [-0.02, 0.02
Subtotal (95% CI)	103	105	ţ	100.0 %	0.0 [-0.02, 0.02
Total events: 0 (Vitamin D ar	. , , ,	nt)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0$.	. ,				
9 Calcipotriol vs. tazarotene	0			100.0.0/	
Guenther 2000	1/60	1/60		100.0 %	0.0 [-0.05, 0.05
Subtotal (95% CI)	60	60	+	100.0 %	0.0 [-0.05, 0.05]
Total events: I (Vitamin D ar	nalogue), I (Other treatmen	nt)			
		-	I -0.5 0 0.5 I		
		Favours vitamin	D analogue Favours other	treatment	

Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M-	Weight	(Continued) Risk Difference M-	
	n/N	n/N	H,Random,95% Cl		H,Random,959 Cl	
Heterogeneity: not applicab						
Test for overall effect: $Z = 0$	0.0 (P = 1.0)					
10 Calcipotriol vs. vitamin B	12 cream					
Stuecker 2001	0/13	0/13		100.0 %	0.0 [-0.14, 0.14]	
Subtotal (95% CI)	13	13	+	100.0 %	0.0 [-0.14, 0.14]	
Total events: 0 (Vitamin D a	nalogue), 0 (Other treatmer	nt)				
Heterogeneity: not applicab	le					
Test for overall effect: $Z = 0$	0.0 (P = 1.0)					
II Head-to-head vitamin D	alone or in combination: tw	rice daily vs OD				
Baiocchi 1997	1/132	1/132	•	25.4 %	0.0 [-0.02, 0.02]	
Guenther 2002 (H)	1/235	0/151	•	59.8 %	0.00 [-0.01, 0.02]	
Kragballe 1998b	3/174	3/174		14.8 %	0.0 [-0.03, 0.03]	
Subtotal (95% CI)	541	457		100.0 %	0.00 [-0.01, 0.01]	
Total events: 5 (Vitamin D a	nalogue), 4 (Other treatmer	nt)				
Heterogeneity: $Tau^2 = 0.0;$	$Chi^2 = 0.18$, $df = 2$ (P = 0.9	I); I ² =0.0%				
Test for overall effect: $Z = 0$	0.47 (P = 0.64)					
12 Head-to-head vitamin D	alone or in combination: no	occlusion vs. occlusion				
Subtotal (95% CI)	0	0			Not estimable	
Total events: 0 (Vitamin D a	nalogue), 0 (Other treatmer	nt)				
Heterogeneity: not applicab	le					
Test for overall effect: not ap	oplicable					

Favours vitamin D analogue Favours other treatment

Analysis 15.9. Comparison 15 Vitamin D analogues versus other treatment, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D analogue Other treatment		Risk Difference M-	Weight	Risk Difference M-	
	n/N	n/N	H,Random,95% Cl		H,Random,9 CI	
I Calcipotriol vs. coal tar						
Alora-Palli 2010	4/30	5/30		19.1 %	-0.03 [-0.21, 0.15]	
Tham 1994	1/30	0/30		80.9 %	0.03 [-0.05, 0.12]	
Subtotal (95% CI)	60	60	+	100.0 %	0.02 [-0.06, 0.10]	
Total events: 5 (Vitamin D an Heterogeneity: Tau ² = 0.0; Cl Test for overall effect: $Z = 0.5$	$hi^2 = 0.69, df = 1 (P = 0.4)$ 51 (P = 0.61)	,				
2 Calcipotriol vs. coal tar poly Pinheiro 1997	ytherapy 15/65	10/57		100.0 %	0.06 [-0.09, 0.20]	
Subtotal (95% CI)	65	57		100.0 %	0.06 [-0.09, 0.20]	
 Total events: 15 (Vitamin D a Heterogeneity: not applicable Test for overall effect: Z = 0.7 3 Calcipotriol vs. nicotinamide Levine 2010 (H) 	e 76 (P = 0.45)	16/48		100.0 %	-0.15 [-0.32, 0.03	
Subtotal (95% CI)	48	48		100.0 %	-0.15 [-0.32, 0.03]	
Total events: 9 (Vitamin D an Heterogeneity: not applicable Test for overall effect: Z = 1.6 4 Calcipotriol vs. calcipotriol Levine 2010 (H)	e 65 (P = 0.099)		ily 	100.0 %	-0.17 [-0.30, -0.03]	
Subtotal (95% CI)	48	144	-	100.0 %	-0.17 [-0.30, -0.03]	
Total events: 9 (Vitamin D an Heterogeneity: not applicable Test for overall effect: Z = 2.4 5 Calcipotriol vs. corticosterc	alogue), 51 (Other treatm e 42 (P = 0.016)					
Crosti 1997	7/80	0/80		100.0 %	0.09 [0.02, 0.15]	
Subtotal (95% CI)	80 halogue), 0 (Other treatme	80	•	100.0 %	0.09 [0.02, 0.15]	

(... Continued)

Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference	Weight	(Continued Risk Difference
	n/N	n/N	M- H,Random,95% Cl		M- H,Random,S CI
Calcipotriol vs. propylthic	ouracil cream				
Sanchez 2001	0/14	1/14		100.0 %	-0.07 [-0.25, 0.11]
Subtotal (95% CI)	14	14	-	100.0 %	-0.07 [-0.25, 0.11]
	analogue), I (Other treatme	nt)			
Heterogeneity: not applicat	ble				
Test for overall effect: $Z = 0$	0.79 (P = 0.43)				
7 Calcipotriol vs. tacrolimus	s ointment				
Ortonne 2006	3/40	43/84		100.0 %	-0.19 [-0.37, -0.01]
Subtotal (95% CI)	40	84	-	100.0 %	-0.19 [-0.37, -0.01]
otal events: 13 (Vitamin D	analogue), 43 (Other treat	ment)			
Heterogeneity: not applicat	ble				
Test for overall effect: $Z = 2$	2.03 (P = 0.042)				
8 Calcipotriol vs. tazaroten	e				
Han 2001	10/101	13/103		100.0 %	-0.03 [-0.11, 0.06]
Subtotal (95% CI)	101	103	•	100.0 %	-0.03 [-0.11, 0.06]
	analogue), 13 (Other treat	ment)			
Heterogeneity: not applicat	ble				
Test for overall effect: Z = (0.62 (P = 0.54)				
Calcipotriol vs. tazaroten	e gel plus mometasone furo	ate cream			
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D a	analogue), 0 (Other treatme	nt)			
Heterogeneity: not applicat	ble				
Fest for overall effect: not a	pplicable				
10 Calcipotriol vs. vitamin E	312 cream				
Stuecker 2001	4/13	1/13		100.0 %	0.23 [-0.06, 0.52]
Subtotal (95% CI)	13	13		100.0 %	0.23 [-0.06, 0.52]
	analogue), I (Other treatme			10000 /0	0.25 [0.000, 0.02]
Heterogeneity: not applicat	e , , ,				
Test for overall effect: Z =					
	, , ,				
Guenther 2002 (H)) alone or in combination: tv 25/235	vice daily vs OD	_	71.9 %	0.01 [-0.05, 0.07]
			J		
Kragballe 1998b	54/173	59/172		28.1 %	-0.03 [-0.13, 0.07]
Subtotal (95% CI)	408	323	+	100.0 %	0.00 [-0.06, 0.05]
	analogue), 74 (Other treat				
- ,	$Chi^2 = 0.49, df = 1 (P = 0.49)$	18); I ² =0.0%			
Test for overall effect: $Z = 0$	0.13 (P = 0.89)				
2 Head-to-head vitamin D) alone or in combination: n	o occlusion vs. occlusion			
Subtotal (95% CI)	0	0			Not estimable
līotal events: 0 (Vitamin D a	analogue), 0 (Other treatme	nt)			
			<u> </u>		
		-C	0.5 -0.25 0 0.25 0.5	5	
		Favours vitamir	n D analogue Favours other	r treatment	
					(Continued

						(Continued)
Study or subgroup	Vitamin D analogue	Other treatment	atment Differe		Weight	Risk Difference M-
	n/N	n/N	H,Ran	idom,95% Cl		H,Random,95% Cl
Heterogeneity: not applica	able					
Test for overall effect: not	applicable					
			<u> </u>		I	
		-C).5 -0.25 (0.25	0.5	
		Favours vitamir	n D analogue	Favours	other treatment	

Analysis 15.10. Comparison 15 Vitamin D analogues versus other treatment, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 15 Vitamin D analogues versus other treatment

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,959 Cl
I Calcipotriol vs. coal tar					
Tham 1994	1/30	1/30	+	100.0 %	0.0 [-0.09, 0.09]
Subtotal (95% CI)	30	30	+	100.0 %	0.0 [-0.09, 0.09]
Total events: I (Vitamin D an	alogue), I (Other treatmen	t)			
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
2 Calcipotriol vs. coal tar poly	ytherapy				
Van de Kerkhof 2002a	0/41	0/47	•	100.0 %	0.0 [-0.04, 0.04]
Subtotal (95% CI)	41	47	•	100.0 %	0.0 [-0.04, 0.04]
Total events: 0 (Vitamin D an	alogue), 0 (Other treatmen	t)			
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
3 Calcipotriol vs. nicotinamid	le 1.4%, twice daily				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D an	alogue), 0 (Other treatmen	t)			
Heterogeneity: not applicable	2				
		-2			
		Favours vitamin	D analogue Favours other	reatment	(Continued)

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup Vita	imin D analogue	Other treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random, C
est for overall effect: not applicable					
Calcipotriol vs. calcipotriol + nicot Subtotal (95% CI)	inamide 1.4%, twice 0	daily 0			Not estimable
otal events: 0 (Vitamin D analogue) Heterogeneity: not applicable Test for overall effect: not applicable is Calcipotriol vs. corticosteroid + sa	licylic acid				
Crosti 1997	0/80	0/80		100.0 %	0.0 [-0.02, 0.02]
Subtotal (95% CI) Total events: 0 (Vitamin D analogue) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = Calcipotriol vs. propylthiouracil cress Sanchez 2001	1.0)	80 0/14		100.0 %	0.0 [-0.02, 0.02]
					0.0 [-0.13, 0.13]
Subtotal (95% CI) Total events: 0 (Vitamin D analogue) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = Calcipotriol vs. tacrolimus ointmer	I.O) ht			100.0 %	0.0 [-0.13, 0.13]
Ortonne 2006	0/40	0/84		100.0 %	0.0 [-0.04, 0.04]
Subtotal (95% CI) Fotal events: 0 (Vitamin D analogue) Heterogeneity: not applicable Fest for overall effect: Z = 0.0 (P = 8 Calcipotriol vs. tazarotene		84		100.0 %	0.0 [-0.04, 0.04]
Han 2001	1/92	2/91	•	100.0 %	-0.01 [-0.05, 0.03
Subtotal (95% CI) Total events: I (Vitamin D analogue) Heterogeneity: not applicable Test for overall effect: Z = 0.59 (P = Calcipotriol vs. tazarotene gel plus	0.55)			100.0 %	-0.01 [-0.05, 0.03
Subtotal (95% CI) Total events: 0 (Vitamin D analogue) Heterogeneity: not applicable Test for overall effect: not applicable		0 nt)			Not estimable
0 Calcipotriol vs. vitamin B12 crear	n				
Subtotal (95% CI) Total events: 0 (Vitamin D analogue) Heterogeneity: not applicable Test for overall effect: not applicable	0), 0 (Other treatmer	0			Not estimable
I Head-to-head vitamin D alone o	r in combination: tw	ice daily vs OD			

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Study or subgroup	Vitamin D analogue	Other treatment	Risk Difference M-	Weight	Risk Difference M-	
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl	
Baiocchi 1997	0/132	0/132		100.0 %	0.0 [-0.01, 0.01]	
Subtotal (95% CI)	132	132		100.0 %	0.0 [-0.01, 0.01]	
Total events: 0 (Vitamin D a	nalogue), 0 (Other treatme	nt)				
Heterogeneity: not applicab	le					
Test for overall effect: $Z = C$	0.0 (P = 1.0)					
12 Head-to-head vitamin D	alone or in combination: no	occlusion vs. occlusion				
Bourke 1993b	0/19	0/19	-	100.0 %	0.0 [-0.10, 0.10]	
Subtotal (95% CI)	19	19	+	100.0 %	0.0 [-0.10, 0.10]	
Total events: 0 (Vitamin D a	nalogue), 0 (Other treatme	nt)				
Heterogeneity: not applicab	le					
Test for overall effect: $Z = C$	0.0 (P = 1.0)					
			-2 -1 0 1 2			

Favours vitamin D analogue Favours other treatment

Analysis 16.1. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 1 IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: I IAGI

Study or subgroup	Active treatment		Placebo		Std. Mean Difference			Std. Mean Difference		
N		Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI				IV,Random,95% CI	
l Betamethasone vale	rate 0.1%, OD									
2 Calcipotriol ointmen	nt, OD									
3 Pimecrolimus cream	, 1% OD/twice daily									
Gribetz 2004	22	-2.95 (1.33)	25	-1.56 (1.23)			+			-1.07 [-1.69, -0.45]
4 Tacrolimus ointment	t 0.1%, twice daily					1		I		
					-10	-5	0	5	10	
				Favou	rs active tr	eatment		Favour	s placebo	

Topical treatments for chronic plaque psoriasis (Review)

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Analysis 16.2. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: I 6 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 2 TSS

Study or subgroup	Active treatment		Placebo		Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% CI
l Betamethasone vale	erate 0.1%, OD					
2 Calcipotriol ointmer	nt, OD					
3 Pimecrolimus cream	n, 1% OD/twice daily					
Gribetz 2004	28	-4.1 (1.03)	29	-2.67 (1.03)	+	-1.37 [-1.95, -0.79]
4 Tacrolimus ointment	t 0.1%, twice daily					
					-10 -5 0 5 1	0
				Favour	s active treatment Favours place	ebo

Analysis 16.3. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 3 PASI

Study or subgroup	Active treatment		Placebo		Std. Mean Difference	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI	IV,Random,95% CI
I Betamethasone valer	rate 0.1%, OD					
Kreuter 2006 (P)	17	2.94 (3.76)	19	13.84 (3.76)		-2.83 [-3.79, -1.88]
2 Calcipotriol ointment	t, OD					
Kreuter 2006 (P)	19	9.68 (3.76)	19	13.84 (3.76)		-1.08 [-1.77, -0.40]
3 Pimecrolimus cream,	1% OD/twice daily					
Kreuter 2006 (P)	20	11.45 (3.76)	19	13.84 (3.76)	+	-0.62 [-1.27, 0.02]
4 Tacrolimus ointment	0.1%, twice daily					
					-10 -5 0 5	10
				Favours	s active treatment Favours	placebo

Analysis 16.4. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 4 PAGI

Study or subgroup	Active treatment		Placebo			Sto Mea Differenc	n	Weight	Std. Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	IV	Random,959	% Cl		IV,Random,95% CI
I Betamethasone valerat	te 0.1%, OD								
Subtotal (95% CI)) 0		0						Not estimable
Heterogeneity: not appli	cable								
Test for overall effect: no	et applicable								
2 Calcipotriol ointment,	OD								
Subtotal (95% CI)) 0		0						Not estimable
Heterogeneity: not appli	cable								
Test for overall effect: no	ot applicable								
3 Pimecrolimus cream, I	% OD/twice daily								
Gribetz 2004	22	-2.36 (0.85)	25	-1.72 (1.06)		-		100.0 %	-0.65 [-1.24, -0.06]
Subtotal (95% CI)	22		25			•		100.0 %	-0.65 [-1.24, -0.06]
Heterogeneity: not appli	cable								
Test for overall effect: Z	= 2.16 (P = 0.031)								
4 Tacrolimus ointment 0	.1%, twice daily								
Subtotal (95% CI)) 0		0						Not estimable
Heterogeneity: not appli	cable								
Test for overall effect: no	ot applicable								
Test for subgroup differe	nces: Not applicable								
					-10 -5	0	5 I	C	

Analysis 16.5. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Active treatment N	Mean(SD)	Placebo N	Mean(SD)		Std. Mean rence n,95% Cl	Weight	Std. Mean Difference IV,Random,95% Cl
I Betamethasone valerate	e 0.1%, OD							
Kreuter 2006 (P)	17	2.94 (3.76)	19	13.84 (3.76)	-		100.0 %	-2.83 [-3.79, -1.88]
Subtotal (95% CI)	17		19		•		100.0 %	-2.83 [-3.79, -1.88]
Heterogeneity: not applic Test for overall effect: Z = 2 Calcipotriol ointment, C	= 5.83 (P < 0.00001) DD							
Kreuter 2006 (P)	19	9.68 (3.76)	19	13.84 (3.76)			100.0 %	-1.08 [-1.77, -0.40]
Subtotal (95% CI) Heterogeneity: not applic Test for overall effect: Z = 3 Pimecrolimus cream, 19	cable = 3.09 (P = 0.0020)		19		•		100.0 %	-1.08 [-1.77, -0.40]
Gribetz 2004	22	-2.95 (1.33)	25	-1.56 (1.23)			52.3 %	-1.07 [-1.69, -0.45]
Kreuter 2006 (P)	20	11.45 (3.76)	19	13.84 (3.76)	-		47.7 %	-0.62 [-1.27, 0.02]
Subtotal (95% CI) Heterogeneity: Tau ² = 0.1 Test for overall effect: Z =	0; Chi ² = 0.97, df = 1 = 3.77 (P = 0.00016)	(P = 0.33); I ² =	44 0.0%		•		100.0 %	-0.86 [-1.30, -0.41]
4 Tacrolimus ointment 0. Subtotal (95% CI)	. ,		0					Not estimable
Heterogeneity: not applic Test for overall effect: not	cable t applicable							ivot estimable
Test for subgroup differer	nces: Chi ² = 13.67, df	T = 2 (P = 0.00),	l ² =85%					
					10 -5 0 ive treatment	5 I (Favours place		

Analysis 16.6. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 6 Total withdrawals

Study or subgroup	Active treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
Betamethasone valerate 0.15	%, OD				
Kreuter 2006 (P)	3/20	1/20	-	100.0 %	0.10 [-0.08, 0.28]
Subtotal (95% CI)	20	20	-	100.0 %	0.10 [-0.08, 0.28]
Total events: 3 (Active treatme	ent), I (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 1.0^{\circ}$	7 (P = 0.29)				
2 Calcipotriol ointment, OD					
Kreuter 2006 (P)	1/20	1/20		100.0 %	0.0 [-0.14, 0.14]
Subtotal (95% CI)	20	20	+	100.0 %	0.0 [-0.14, 0.14]
Total events: I (Active treatme	ent), I (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
3 Pimecrolimus cream, 1% OE	D/twice daily				
Gribetz 2004	2/28	4/29	-	39.7 %	-0.07 [-0.22, 0.09]
Kreuter 2006 (P)	0/20	1/20	-	60.3 %	-0.05 [-0.18, 0.08]
Subtotal (95% CI)	48	49	•	100.0 %	-0.06 [-0.16, 0.04]
Total events: 2 (Active treatme	ent), 5 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$; Ch	$hi^2 = 0.03$, $df = 1$ (P = 0.8	6); l ² =0.0%			
Test for overall effect: $Z = 1.13$	2 (P = 0.26)				
4 Tacrolimus ointment 0.1%, tv	wice daily				
Lebwohl 2004	4/ 2	16/55		100.0 %	-0.17 [-0.30, -0.03]
Subtotal (95% CI)	112	55	•	100.0 %	-0.17 [-0.30, -0.03]
Total events: 14 (Active treatm	nent), 16 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 2.4$	I (P = 0.016)				
T	$Chi^2 = 5.96, df = 3 (P = 0)$	$(11) 1^2 = 50\%$			

Analysis 16.7. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Active treatment	Placebo	Risk Difference M-	Weight	Risk Difference M
	n/N	n/N	H,Random,95% Cl		M- H,Random,9 CI
Betamethasone valerate 0	.1%, OD				
Kreuter 2006 (P)	0/20	0/20		100.0 %	0.0 [-0.09, 0.09]
Subtotal (95% CI)	20	20	+	100.0 %	0.0 [-0.09, 0.09]
Total events: 0 (Active treat	ment), 0 (Placebo)				
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 0$.0 (P = 1.0)				
2 Calcipotriol ointment, OD)				
Kreuter 2006 (P)	0/20	0/20	-	100.0 %	0.0 [-0.09, 0.09]
Subtotal (95% CI)	20	20	+	100.0 %	0.0 [-0.09, 0.09]
Total events: 0 (Active treat	ment), 0 (Placebo)				
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 0$.0 (P = 1.0)				
3 Pimecrolimus cream, 1% C	DD/twice daily				
Gribetz 2004	0/28	0/29	-	66.2 %	0.0 [-0.07, 0.07]
Kreuter 2006 (P)	0/20	0/20	+	33.8 %	0.0 [-0.09, 0.09]
Subtotal (95% CI)	48	49	•	100.0 %	0.0 [-0.05, 0.05]
Total events: 0 (Active treat	ment), 0 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$; ($Chi^2 = 0.0, df = 1 (P = 1.00)$); I ² =0.0%			
Test for overall effect: $Z = 0$.0 (P = 1.0)				
4 Tacrolimus ointment 0.1%	twice daily				
Lebwohl 2004	0/112	1/55	-	100.0 %	-0.02 [-0.06, 0.03]
Subtotal (95% CI)	112	55	•	100.0 %	-0.02 [-0.06, 0.03]
Total events: 0 (Active treat	ment), I (Placebo)				
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 0$.81 (P = 0.42)				
Test for subgroup difference	s: $Chi^2 = 0.35$, $df = 3$ (P = 0)	0.95), I ² =0.0%			

-I -0.5 0 0.5 I

Analysis 16.8. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Active treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Betamethasone valerate 0	.1%, OD				
Kreuter 2006 (P)	0/20	1/20	=	100.0 %	-0.05 [-0.18, 0.08]
Subtotal (95% CI)	20	20	•	100.0 %	-0.05 [-0.18, 0.08]
Total events: 0 (Active treat	ment), I (Placebo)				
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 0$	0.77 (P = 0.44)				
2 Calcipotriol ointment, OD)				
Kreuter 2006 (P)	0/20	1/20		100.0 %	-0.05 [-0.18, 0.08]
Subtotal (95% CI)	20	20	•	100.0 %	-0.05 [-0.18, 0.08]
Total events: 0 (Active treat	ment), I (Placebo)				
Heterogeneity: not applicabl	le				
Test for overall effect: $Z = 0$	0.77 (P = 0.44)				
3 Pimecrolimus cream, 1% (DD/twice daily				
Gribetz 2004	1/28	2/29	+	55.3 %	-0.03 [-0.15, 0.08]
Kreuter 2006 (P)	0/20	1/20	-	44.7 %	-0.05 [-0.18, 0.08]
Subtotal (95% CI)	48	49	•	100.0 %	-0.04 [-0.13, 0.04]
Total events: (Active treat	ment), 3 (Placebo)				
Heterogeneity: $Tau^2 = 0.0;$	$Chi^2 = 0.04, df = 1 (P = 0.8)$	5); I ² =0.0%			
Test for overall effect: $Z = 0$	0.93 (P = 0.35)				
4 Tacrolimus ointment 0.1%	, twice daily				
Lebwohl 2004	0/112	6/55	-	100.0 %	-0.11 [-0.19, -0.02]
Subtotal (95% CI)	112	55	•	100.0 %	-0.11 [-0.19, -0.02]
Total events: 0 (Active treat	ment), 6 (Placebo)				
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 2$.52 (P = 0.012)				
Test for subgroup difference	e^{-2} Chi ² = 1.44 df = 3 (P =	(0.70) $I^2 = 0.0\%$			

-I -0.5 0 0.5 I

Analysis 16.9. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 9 Adverse events (local)

Study or subgroup	Active treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
Betamethasone valerate 0.19	%, OD				
Kreuter 2006 (P)	0/20	1/20		100.0 %	-0.05 [-0.18, 0.08]
Subtotal (95% CI)	20	20	•	100.0 %	-0.05 [-0.18, 0.08]
Total events: 0 (Active treatme	ent), I (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.77$	7 (P = 0.44)				
2 Calcipotriol ointment, OD					
Kreuter 2006 (P)	2/20	1/20		100.0 %	0.05 [-0.11, 0.21]
Subtotal (95% CI)	20	20	•	100.0 %	0.05 [-0.11, 0.21]
Total events: 2 (Active treatme	ent), I (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.60$	0 (P = 0.55)				
3 Pimecrolimus cream 1%, OE)/twice daily				
Gribetz 2004	1/28	1/29	-	58.3 %	0.00 [-0.09, 0.10]
Kreuter 2006 (P)	5/20	1/20	-	41.7 %	0.20 [-0.01, 0.41]
Subtotal (95% CI)	48	49	+	100.0 %	0.08 [-0.15, 0.31]
Total events: 6 (Active treatme	ent), 2 (Placebo)				
Heterogeneity: $Tau^2 = 0.02$; C	$2hi^2 = 3.99, df = 1 (P = 0.9)$	05); I ² =75%			
Test for overall effect: $Z = 0.72$	2 (P = 0.47)				
4 Tacrolimus ointment 0.1%, tv	wice daily				
Lebwohl 2004	4/ 2	16/55		100.0 %	-0.17 [-0.30, -0.03]
Subtotal (95% CI)	112	55	•	100.0 %	-0.17 [-0.30, -0.03]
Total events: 14 (Active treatm	nent), 16 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 2.4$,				
T . C . I	$Chi^2 = 5.61, df = 3 (P = 0)$	$0, 3), ^2 = 47\%$			

Analysis 16.10. Comparison 16 Flexural/facial psoriasis: placebo-controlled trials, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 16 Flexural/facial psoriasis: placebo-controlled trials

Outcome: 10 Adverse events (systemic)

Study or subgroup	Active treatment	Placebo	Risk Difference M- H.Random,95%	Risk Difference M- H.Random,95%	
	n/N	n/N	Cl	CI	
l Betamethasone valerate ().1%, OD				
2 Calcipotriol ointment, OI	C				
3 Pimecrolimus cream 1%,	OD/twice daily				
4 Tacrolimus ointment 0.1%	6, twice daily				
Lebwohl 2004	0/112	0/55	+	0.0 [-0.03, 0.03]	
			-0.5 -0.25 0 0.25 0.5		
		Fav	ours active treatment Favours placebo		

Analysis 17.1. Comparison 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment, Outcome I IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

Outcome: I IAGI

Study or subgroup	Vitamin D alone or in combination		Other treatment			Std. Iean ence	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,	95% CI		IV,Random,95% CI
I Calcipotriol vs. BMV								
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicable	e							
Test for overall effect: not ap	plicable							
2 Calcipotriol vs. calcipotriol	+ hydrocortison	e						
Ortonne 2010	202	1.37 (1.16)	206	1.04 (1.02)	+		100.0 %	0.30 [0.11, 0.50]
Subtotal (95% CI)	202		206		•	:	100.0 %	0.30 [0.11, 0.50]
Heterogeneity: not applicable	e							
Test for overall effect: $Z = 3$.	.03 (P = 0.0024)							
3 Calcipotriol vs. calcitriol								
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicable	e							
Test for overall effect: not ap	plicable							
4 Calcipotriol vs. pimecrolim	nus							
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicable	e							
Test for overall effect: not ap	plicable							
5 Calcitriol vs. tacrolimus								
Liao 2007	24	-3.67 (1.78)	25	-4.48 (2.02)		-	100.0 %	0.42 [-0.15, 0.98]
Subtotal (95% CI)	24		25		•	. :	100.0 %	0.42 [-0.15, 0.98]
Heterogeneity: not applicable	e							
Test for overall effect: $Z = 1$.								
Test for subgroup differences	· ,	= I (P = 0.70), l² =0.0%					
				-4	-2 0	2 4		
			Favours vita	min D alone or in o	combination	Favours other ti	reatment	

Analysis 17.2. Comparison 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

Outcome: 2 TSS

Vr Study or subgroup	tamin D alone or in combination	C	Other treatment		C	Std. Mean Vifference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Ran	dom,95% Cl		IV,Random,95% CI
I Calcipotriol vs. BMV								
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicable								
Test for overall effect: not appl	licable							
2 Calcipotriol vs. calcipotriol +	- hydrocortisone							
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicable								
Test for overall effect: not appl	licable							
3 Calcipotriol vs. calcitriol								
Ortonne 2003	75	1.78 (1.5)	75	0.86 (1.5)			100.0 %	0.61 [0.28, 0.94]
Subtotal (95% CI)	75		75			•	100.0 %	0.61 [0.28, 0.94]
Heterogeneity: not applicable								
Test for overall effect: $Z = 3.65$	5 (P = 0.00026)							
4 Calcipotriol vs. pimecrolimus	5							
Subtotal (95% CI)	0		0					Not estimable
Heterogeneity: not applicable								
Test for overall effect: not appl	licable							
5 Calcitriol vs. tacrolimus								
Liao 2007	24	-0.51 (0.35)	25	-0.64 (0.51)		H	100.0 %	0.29 [-0.27, 0.85]
Subtotal (95% CI)	24		25			•	100.0 %	0.29 [-0.27, 0.85]
Heterogeneity: not applicable								
Test for overall effect: $Z = 1.0$	I (P = 0.3I)							
Test for subgroup differences:	Chi ² = 0.92, df =	= (P = 0.34), l ²	=0.0%					
							1	
				-4	-2	0 2	4	
			Favours vitar	nin D alone or in co	mbination	Favours oth	ner treatment	

Analysis 17.3. Comparison 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment, Outcome 3 PASI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

Outcome: 3 PASI

			Other treatment		Mean Difference	Weight	Mean Difference
	Ν	Mean(SD)	N	Mean(SD)	IV,Random,95% Cl		IV,Random,95% C
Calcipotriol vs. BMV							
Kreuter 2006 (H)	19	9.68 (3.26)	17	2.94 (3.26)		100.0 %	2.02 [1.20, 2.84]
ubtotal (95% CI)	19		17		•	100.0 %	2.02 [1.20, 2.84]
eterogeneity: not applicable							
est for overall effect: Z = 4.83 (P	< 0.00001)						
Calcipotriol vs. calcipotriol + hyd	drocortison	e					
Ortonne 2010	202	0.21 (0.3)	206	0.13 (0.19)	-+-	100.0 %	0.32 [0.12, 0.51]
ubtotal (95% CI)	202		206		•	100.0 %	0.32 [0.12, 0.51]
eterogeneity: not applicable							
est for overall effect: Z = 3.20 (P	= 0.0014)						
Calcipotriol vs. calcitriol							
ubtotal (95% CI)	0		0				Not estimable
eterogeneity: not applicable							
est for overall effect: not applicab	le						
Calcipotriol vs. pimecrolimus							
Kreuter 2006 (H)	19	9.68 (3.26)	20	11.45 (3.26)		100.0 %	-0.53 [-1.17, 0.11]
ubtotal (95% CI)	19		20		•	100.0 %	-0.53 [-1.17, 0.11]
eterogeneity: not applicable							
est for overall effect: $Z = 1.63$ (P	= 0.10)						
Calcitriol vs. tacrolimus							
ubtotal (95% CI)	0		0				Not estimable
eterogeneity: not applicable							
est for overall effect: not applicab							
est for subgroup differences: Chi ²	= 23.31, d	f = 2 (P = 0.0)	10), I ² =91%				

Analysis 17.5. Comparison 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

V Study or subgroup	/itamin D alone or in combination N	Mean(SD)	Other treatment N	Mean(SD)	Std. Mean Difference IV.Random.95% Cl	Weight	Std. Mean Difference IV.Random,95% CI
	14	1 Icaii(3D)	14	r icar(JD)			IV, Randol II, 7576 CI
I Calcipotriol vs. BMV						_	
Kreuter 2006 (H)	19	9.68 (3.26)	17	2.94 (3.26)		100.0 %	2.02 [1.20, 2.84]
Subtotal (95% CI)	19		17			100.0 %	2.02 [1.20, 2.84]
Heterogeneity: not applicable	2						
Test for overall effect: $Z = 4.4$	83 (P < 0.00001))					
2 Calcipotriol vs. calcipotriol	+ hydrocortison	e					
Ortonne 2010	202	1.37 (1.16)	206	1.04 (1.02)		100.0 %	0.30 [0.11, 0.50]
Subtotal (95% CI)	202		206		•	100.0 %	0.30 [0.11, 0.50]
Heterogeneity: not applicable	2						
Test for overall effect: $Z = 3.0$	03 (P = 0.0024)						
3 Calcipotriol vs. calcitriol							
Ortonne 2003	75	1.78 (1.5)	75	0.86 (1.5)		100.0 %	0.61 [0.28, 0.94]
Subtotal (95% CI)	75		75		•	100.0 %	0.61 [0.28, 0.94]
Heterogeneity: not applicable	2						
Test for overall effect: $Z = 3.6$	65 (P = 0.00026))					
4 Calcipotriol vs. pimecrolim	US						
Kreuter 2006 (H)	19	9.68 (3.26)	20	11.45 (3.26)		100.0 %	-0.53 [-1.17, 0.11]
Subtotal (95% CI)	19		20		-	100.0 %	-0.53 [-1.17, 0.11]
Heterogeneity: not applicable	2						
Test for overall effect: $Z = 1.6$	63 (P = 0.10)						
5 Calcitriol vs. tacrolimus							
Liao 2007	24	-3.67 (1.78)	25	-4.48 (2.02)	+	100.0 %	0.42 [-0.15, 0.98]
Subtotal (95% CI)	24		25		-	100.0 %	0.42 [-0.15, 0.98]
Heterogeneity: not applicable	2						
Test for overall effect: $Z = 1$.	45 (P = 0.15)						
Test for subgroup differences	: Chi ² = 25.69, d	If = 4 (P = 0.0)	00), l ² =84%				
						1	

Favours vitamin D alone or in combination

Favours other treatment

Analysis 17.6. Comparison 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

Outcome: 6 Total withdrawals

	Vitamin D alone or in		Risk		Risk
Study or subgroup	combination	Other treatment	Difference M-	Weight	Difference M
	n/N	n/N	H,Random,95% Cl		H,Random, C
I Calcipotriol vs. BMV					
Kreuter 2006 (H)	1/20	3/20		100.0 %	-0.10 [-0.28, 0.08]
Subtotal (95% CI)	20	20	-	100.0 %	-0.10 [-0.28, 0.08]
Total events: I (Vitamin D ale	one or in combination), 3	8 (Other treatment)			
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 1$.	07 (P = 0.29)				
2 Calcipotriol vs. calcipotriol	+ hydrocortisone				
Ortonne 2010	34/202	27/206		100.0 %	0.04 [-0.03, 0.11]
Subtotal (95% CI)	202	206	•	100.0 %	0.04 [-0.03, 0.11]
Total events: 34 (Vitamin D a	lone or in combination),	27 (Other treatment)			
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 1$.	06 (P = 0.29)				
3 Calcipotriol vs. calcitriol					
Ortonne 2003	10/75	10/75		100.0 %	0.0 [-0.11, 0.11
Subtotal (95% CI)	75	75	+	100.0 %	0.0 [-0.11, 0.11
Total events: 10 (Vitamin D a	lone or in combination),	10 (Other treatment)			
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.1$	0 (P = 1.0)				
4 Calcipotriol vs. pimecrolim	us				
Kreuter 2006 (H)	1/20	0/20		100.0 %	0.05 [-0.08, 0.18
Subtotal (95% CI)	20	20	•	100.0 %	0.05 [-0.08, 0.18]
Total events: I (Vitamin D ale	one or in combination), () (Other treatment)			
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.7$	77 (P = 0.44)				
5 Calcitriol vs. tacrolimus					
Liao 2007	3/25	0/25		100.0 %	0.12 [-0.02, 0.26
Subtotal (95% CI)	25	25	•	100.0 %	0.12 [-0.02, 0.26]
Total events: 3 (Vitamin D ale	one or in combination), () (Other treatment)			
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 1$.	66 (P = 0.096)				
	: Chi ² = 3.91, df = 4 (P	= 0.42), I ² =0.0%			
Test for subgroup differences					

Analysis 17.7. Comparison 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

Outcome: 7 Withdrawals due to adverse events

	Vitamin D alone or in		Risk		Risk
Study or subgroup	combination	Other treatment	Difference M-	Weight	Difference
	n/N	n/N	H,Random,95% Cl		H,Random, C
I Calcipotriol vs. BMV					
Kreuter 2006 (H)	0/20	0/20	-	100.0 %	0.0 [-0.09, 0.09
Subtotal (95% CI)	20	20	+	100.0 %	0.0 [-0.09, 0.09]
Total events: 0 (Vitamin D alc	ne or in combination), 0	(Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$) (P = 1.0)				
2 Calcipotriol vs. calcipotriol	+ hydrocortisone				
Ortonne 2010	19/202	6/206		100.0 %	0.06 [0.02, 0.11]
Subtotal (95% CI)	202	206	•	100.0 %	0.06 [0.02, 0.11]
Total events: 19 (Vitamin D a	one or in combination),	6 (Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 2.7$	75 (P = 0.0060)				
3 Calcipotriol vs. calcitriol					
Ortonne 2003	9/75	2/75		100.0 %	0.09 [0.01, 0.18
Subtotal (95% CI)	75	75	•	100.0 %	0.09 [0.01, 0.18
Total events: 9 (Vitamin D alc	ne or in combination), 2	(Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 2.2$	23 (P = 0.026)				
4 Calcipotriol vs. pimecrolimu	IS				
Kreuter 2006 (H)	0/20	0/20		100.0 %	0.0 [-0.09, 0.09
Subtotal (95% CI)	20	20	+	100.0 %	0.0 [-0.09, 0.09]
Total events: 0 (Vitamin D alc	ne or in combination), 0	(Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$) (P = 1.0)				
5 Calcitriol vs. tacrolimus					
Liao 2007	0/25	0/25		100.0 %	0.0 [-0.07, 0.07
Subtotal (95% CI)	25	25	+	100.0 %	0.0 [-0.07, 0.07
Total events: 0 (Vitamin D alc	ne or in combination), 0	(Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$) (P = 1.0)				
	$CU^2 = E^2 E^2 = 4 (D - 4)$	$= 0.26$) $l^2 = 24\%$			
Test for subgroup differences:	$Chi^2 = 5.25, df = 4 (P =$	- 0.20), 1 -21/0			

Analysis 17.8. Comparison 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

Outcome: 8 Withdrawals due to treatment failure

	Vitamin D alone or in		Risk		Risk
Study or subgroup	combination	Other treatment	Difference M-	Weight	Difference
	n/N	n/N	H,Random,95% Cl		H,Random, C
I Calcipotriol vs. BMV					
Kreuter 2006 (H)	0/20	0/20	-	100.0 %	0.0 [-0.09, 0.09]
Subtotal (95% CI)	20	20	+	100.0 %	0.0 [-0.09, 0.09]
Total events: 0 (Vitamin D alon	e or in combination), () (Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
2 Calcipotriol vs. calcipotriol +	hydrocortisone				
Ortonne 2010	8/202	4/206		100.0 %	0.02 [-0.01, 0.05]
Subtotal (95% CI)	202	206	•	100.0 %	0.02 [-0.01, 0.05]
Total events: 8 (Vitamin D alon	e or in combination), 4	(Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: Z = 1.20	(P = 0.23)				
3 Calcipotriol vs. calcitriol					
Ortonne 2003	0/75	0/75	-	100.0 %	0.0 [-0.03, 0.03
Subtotal (95% CI)	75	75	+	100.0 %	0.0 [-0.03, 0.03
Total events: 0 (Vitamin D alon	e or in combination), () (Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
4 Calcipotriol vs. pimecrolimus					
Kreuter 2006 (H)	0/20	0/20		100.0 %	0.0 [-0.09, 0.09
Subtotal (95% CI)	20	20	+	100.0 %	0.0 [-0.09, 0.09]
Total events: 0 (Vitamin D alon	e or in combination), () (Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
5 Calcitriol vs. tacrolimus					
Liao 2007	0/25	0/25		100.0 %	0.0 [-0.07, 0.07
Subtotal (95% CI)	25	25	+	100.0 %	0.0 [-0.07, 0.07]
Total events: 0 (Vitamin D alon	e or in combination), () (Other treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
Test for subgroup differences: ($Chi^2 = 0.98, df = 4 (P = 1)$	= 0.9 I), I ² =0.0%			
0 1			1		

Analysis 17.9. Comparison 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 17 Flexural/facial psoriasis: vitamin D alone or in combination versus other treatment

Outcome: 9 Adverse events (local)

	Vitamin D alone or in		Risk		Risk
Study or subgroup	combination	Other treatment	Difference	Weight	Difference
			M- H,Random,95%		M. H,Random,
	n/N	n/N	CI		C
I calcipotriol vs. BMV					
Kreuter 2006 (H)	2/20	0/20	-	100.0 %	0.10 [-0.05, 0.25
Subtotal (95% CI)	20	20	•	100.0 %	0.10 [-0.05, 0.25]
Total events: 2 (Vitamin D al	one or in combination), () (Other treatment)			
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 1$.	28 (P = 0.20)				
2 Calcipotriol vs. calcipotriol	+ hydrocortisone				
Ortonne 2010	52/200	22/204		100.0 %	0.15 [0.08, 0.23]
Subtotal (95% CI)	200	204	•	100.0 %	0.15 [0.08, 0.23]
Total events: 52 (Vitamin D a	alone or in combination),	22 (Other treatment)			
Heterogeneity: not applicable	e				
Test for overall effect: $Z = 4$.	.02 (P = 0.000059)				
3 Calcipotriol vs. calcitriol					
Ortonne 2003	8/75	1/75		100.0 %	0.09 [0.02, 0.17
Subtotal (95% CI)	75	75	•	100.0 %	0.09 [0.02, 0.17
Total events: 8 (Vitamin D al	one or in combination),	l (Other treatment)			
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 2$.	45 (P = 0.014)				
4 Calcipotriol vs. pimecrolim	us				
Kreuter 2006 (H)	2/20	5/20		100.0 %	-0.15 [-0.38, 0.08
Subtotal (95% CI)	20	20	-	100.0 %	-0.15 [-0.38, 0.08]
Total events: 2 (Vitamin D al	one or in combination), 5	ō (Other treatment)			
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 1$.	27 (P = 0.20)				
5 Calcitriol vs. tacrolimus					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D al	one or in combination), () (Other treatment)			
Heterogeneity: not applicabl	e				
Test for overall effect: not ap					
		= 0.10), l ² =53%			

Analysis 18.1. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 1 IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: I IAGI

Std. Mean Difference IV,Random,95% CI	Weight	Std. Mean Difference IV,Random,95% Cl	Mean(SD)	Placebo N	Mean(SD)	Active treatment N	Study or subgroup
							I Vitamin D: calcipotriol
-1.09 [-1.69, -0.48]	37.9 %		-1.25 (1.22)	24	-2.52 (1.08)	25	Green 1994
-0.50 [-0.71, -0.29]	62.1 %		2.41 (1.06)	136	1.85 (1.15)	272	Jemec 2008 (P)
-0.72 [-1.28, -0.16]	100.0 %	•		160		297	Subtotal (95% CI)
				9%	$P = 0.07$); $I^2 = 6$	2.53 (P = 0.011)	Heterogeneity: $Tau^2 = 0.12$; Test for overall effect: $Z = 2$ 2 Potent steroid: betametha
-0.86 [-1.79, 0.06]	4.3 %		-0.9 (1)	10	-1.8 (1)	10	Elie 1983
-1.10 [-1.30, -0.91]	95.7 %	•	2.41 (1.06)	136	1.23 (1.07)	556	Jemec 2008 (P)
-1.09 [-1.29, -0.90]	100.0 %	•		146		566	Subtotal (95% CI)
N 11					= 0.62); I ² =0.0	11.15 (P < 0.00001) asone valerate	Heterogeneity: $Tau^2 = 0.0$; Test for overall effect: $Z = 1$ 3 Potent steroid: betametha
Not estimable				0		pplicable nonide	Subtotal (95% CI) Heterogeneity: not applicab Test for overall effect: not ap 4 Very potent steroid: amoi
-1.42 [-1.80, -1.04]	100.0 %	-	-2.8 (1.23)	67	-4.6 (1.29)	65	Ellis 1988
-1.42 [-1.80, -1.04]	100.0 %	•		67		7.27 (P < 0.00001)	Subtotal (95% CI) Heterogeneity: not applicab Test for overall effect: Z = 7 5 Very potent steroid: clobe
-1.99 [-2.53, -1.45]	20.9 %	-	2.6 (0.955416)	40	41	0.707317 (0.928545)	Cook-Bolden 2010
-1.67 [-1.90, -1.43]	79.1 %		-1.7 (1.09)	189	-3.65 (1.24)	188	Olsen 1991
-1.73 [-1.99, -1.48]	1 00.0 %	•		229 4%	P = 0.28); ² =	I3.22 (P < 0.0000Ⅰ)	Subtotal (95% CI) Heterogeneity: $Tau^2 = 0.01$; Test for overall effect: $Z = 1$ 6 Very potent steroid: halcir
-1.11 [-1.69, -0.53]	100.0 %	+	-1.3 (0.82)	27	-2.3 (0.95)	27	Lepaw 1978
-1.11 [-1.69, -0.53]	100.0 %	•		27			Subtotal (95% CI) Heterogeneity: not applicab Test for overall effect: Z = 3

(Continued . . .)

					Std.		(Continued Std.
Study or subgroup A	0 1		Placebo		Mean Difference	Weight	Mean Difference IV,Random,95% CI
		Mean(SD)		Mean(SD)	IV,Random,95% CI	vveignt	
7 Vitamin D in combination: c	alcipotriol + BMD						,
Jemec 2008 (P)	541	1.09 (1.02)	136	2.41 (1.06)		52.5 %	-1.28 [-1.48, -1.08]
Tyring 2010	135	1.05 (1.15)	42	1.79 (1.28)	-	47.5 %	-0.62 [-0.98, -0.27]
Subtotal (95% CI)	676		178		•	100.0 %	-0.97 [-1.61, -0.32]
Heterogeneity: $Tau^2 = 0.20$; ($Chi^2 = 10.15, df = 1$	$(P = 0.00); ^2 =$	=90%				, [,]
Test for overall effect: $Z = 2.9$	5 (P = 0.0032)						
8 Other treatment: betameth	asone-17,21-dipropi	onate plus salicy	lic acid				
Elie 1983	10	-2.45 (1)	10	-0.9 (1)		100.0 %	-1.48 [-2.50, -0.47]
Subtotal (95% CI)	10		10		•	100.0 %	-1.48 [-2.50, -0.47]
Heterogeneity: not applicable							
Test for overall effect: $Z = 2.8$	6 (P = 0.0042)						
9 Other treatment: ciclopirox	olamine shampoo						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicable							
Test for overall effect: not app	licable						
10 Other treatment: fluocinol	one acetonide, plus	occlusion					
Pauporte 2004	42	-4.38 (0.99)	42	-2.76 (1.57)	-	100.0 %	-1.22 [-1.69, -0.76]
Subtotal (95% CI)	42		42		•	100.0 %	-1.22 [-1.69, -0.76]
Heterogeneity: not applicable							
Test for overall effect: $Z = 5.1$	2 (P < 0.00001)						
I I Other treatment: salicylic a	acid						
Elie 1983	10	-1.8 (1)	10	-0.9 (1)		100.0 %	-0.86 [-1.79, 0.06]
Subtotal (95% CI)	10		10		•	100.0 %	-0.86 [-1.79, 0.06]
Heterogeneity: not applicable							
Test for overall effect: $Z = 1.8$	2 (P = 0.068)						
Test for subgroup differences:	$Chi^2 = 22.17, df = 8$	$B (P = 0.00), I^2 =$	=64%				
				1			
				-10	-5 0 5	10	
				Favours active	treatment Favours pla	acebo	

Analysis 18.2. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: 2 TSS

Difference IV,Random,95% C	Weight	Std. Mean Difference IV,Random,95% CI	Mean(SD)	Placebo N	Mean(SD)	Active treatment N	Study or subgroup
		_					l Vitamin D: calcipotriol
-0.64 [-1.22, -0.07]	11.5 %	_	5.3 (2.5)	24	3.6 (2.7)	25	Green 1994
-0.42 [-0.62, -0.21]	88.5 %	-	4.4 (2.4)	136	3.4 (2.4)	272	Jemec 2008 (P)
-0.44 [-0.64, -0.25]	100.0 %	•		160		297	Subtotal (95% CI)
				0.0%	$(P = 0.47); I^2 =$	4.43 (P < 0.00001)	Heterogeneity: Tau ² = 0.0 Test for overall effect: Z = 2 Potent steroid: betamet
-0.71 [-1.62, 0.20]	4.4 %		3.87 (2.27)	10	2.18 (2.27)	10	Elie 1983
-1.02 [-1.21, -0.82]	95.6 %		4.4 (2.4)	136	2.2 (2.1)	556	Jemec 2008 (P)
-1.00 [-1.19, -0.81]	1 00.0 %	•		146 0.0%	(P = 0.52); I ² =	566 b; Chi ² = 0.41, df = 1	Subtotal (95% CI) Heterogeneity: Tau ² = 0.0
						· · · · · ·	Test for overall effect: Z = 3 Potent steroid: betamet
-1.40 [-1.75, -1.05]	100.0 %		5.97 (2.27)	57	2.78 (2.27)	115	Franz 1999
-1.40 [-1.75, -1.05]	100.0 %	•	-2.7 (2.27)	57 67	-6.31 (2.27)	7.81 (P < 0.00001)	Subtotal (95% CI) Heterogeneity: not applica Test for overall effect: Z = 4 Very potent steroid: am Ellis 1988
-1.58 [-1.98, -1.18]	100.0 %			(=		59	Subtotal (95% CI)
-1.58 [-1.98, -1.18]		•		67			
-1.58 [-1.98, -1.18] -1.58 [-1.98, -1.18]		•		6/		7.70 (P < 0.00001)	Heterogeneity: not applica Test for overall effect: Z = 5 Very potent steroid: cloi
		•	5.4 (2.27)	67	2.1 (2.27)	7.70 (P < 0.00001)	Heterogeneity: not applica Test for overall effect: Z =
-1.58 [-1.98, -1.18] -1.45 [-1.78, -1.11]	100.0 %	•	5.4 (2.27) 5.49 (1.74)		2.1 (2.27) 3.2 (1.74)	= 7.70 (P < 0.00001) betasol propionate	Heterogeneity: not applica Test for overall effect: Z = 5 Very potent steroid: clo
-1.58 [-1.98, -1.18] -1.45 [-1.78, -1.11] -1.31 [-1.69, -0.93]	100.0 % 30.6 %	•	~ /	63	· · ·	: 7.70 (P < 0.00001) betasol propionate 125	Heterogeneity: not applica Test for overall effect: Z = 5 Very potent steroid: clo Franz 2000
-1.58 [-1.98, -1.18]	100.0 % 30.6 % 26.3 % 43.1 %	• • •	5.49 (1.74)	63 47 189 299	3.2 (1.74) 2.4 (2.27)	7.70 (P < 0.00001) betasol propionate 125 95 188 408	Heterogeneity: not applica Test for overall effect: Z = 5 Very potent steroid: cloi Franz 2000 Jarratt 2004 Olsen 1991 Subtotal (95% CI)
-1.58 [-1.98, -1.18] -1.45 [-1.78, -1.11] -1.31 [-1.69, -0.93] -1.71 [-1.95, -1.48]	100.0 % 30.6 % 26.3 % 43.1 %	• • • • •	5.49 (1.74)	63 47 189 299	3.2 (1.74) 2.4 (2.27) 2 (P = 0.16); l ²	7.70 (P < 0.00001) betasol propionate 125 95 188 408 22; Chi ² = 3.73, df = 2 12; Chi ² = 3.73, df = 2	Heterogeneity: not applica Test for overall effect: Z = 5 Very potent steroid: cloi Franz 2000 Jarratt 2004 Olsen 1991

(Continued ...)

Study or subgroup	Active treatment	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	(Continued Std. Mean Difference IV,Random,95% CI
Heterogeneity: not applicab	ble						
Test for overall effect: not a							
7 Vitamin D in combination			127	4.4.(2.4)		F 4 4 0/	
Jemec 2008 (P)	541	1.9 (2.1)	136	4.4 (2.4)		54.4 %	-1.15 [-1.35, -0.96]
Tyring 2010	135	2.1 (2.4)	42	3.7 (2.6)		45.6 %	-0.65 [-1.00, -0.30]
Subtotal (95% CI)	676		178		•	100.0 %	-0.92 [-1.42, -0.43]
Heterogeneity: $Tau^2 = 0.11$ Test for overall effect: $Z = 2$	3.69 (P = 0.00023)	. ,					
8 Other treatment: betame Elie 1983	thasone-17,21-dipro	I.I.5 (2.27)	ICYIIC ACIO	3.87 (2.27)		100.0 %	-1.15 [-2.11, -0.19]
Subtotal (95% CI)	10	~ /	10	~ /	-	100 0 %	-1.15 [-2.11, -0.19]
Heterogeneity: not applicab Test for overall effect: $Z = 2$ 9 Other treatment: ciclopin	ole 2.34 (P = 0.019))	10			100.0 /0	
Shuttleworth 1998	28	4.25 (2.49)	9	4.44 (2.65)		100.0 %	-0.07 [-0.82, 0.68]
Subtotal (95% CI)	28		9		-	100.0 %	-0.07 [-0.82, 0.68]
Heterogeneity: not applicab Test for overall effect: $Z = 0$							
10 Other treatment: fluocir	nolone acetonide, pl	us occlusion					
Pauporte 2004	42	3 (1.71)	42	4.76 (2.18)		100.0 %	-0.89 [-1.34, -0.44]
Subtotal (95% CI) Heterogeneity: not applicab Test for overall effect: Z = 3			42		•	100.0 %	-0.89 [-1.34, -0.44]
II Other treatment: salicyli Elie 1983	c acid 10	2.51 (2.27)	10	3.87 (2.27)		100.0 %	-0.57 [-1.47, 0.32]
Subtotal (95% CI)	10	. /	10	· · ·		100.0 %	-0.57 [-1.47, 0.32]
Heterogeneity: not applicab Test for overall effect: Z =	ble		10			100.0 /0	0.97 [*1.17, 0.9 2]

-2 0 Favours active treatment Favours placebo

2 4

-4

Analysis 18.4. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 4 PAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: 4 PAGI

Stø Mea Differenc IV,Random,95% (Weight	Std. Mean Difference IV,Random,95% Cl	Mean(SD)	Placebo N	Mean(SD)	Active treatment N	Study or subgroup
· ·			()		()		I Vitamin D: calcipotriol
-1.05 [-1.65, -0.45	40.0 %		-1.42 (1.1)	24	-2.52 (0.96)	25	Green 1994
-0.41 [-0.62, -0.20	60.0 %	-	-2.77 (1.8)	135	-3.52 (1.86)	266	Jemec 2008 (P)
-0.66 [-1.28, -0.05	100.0 %	•		159		291	Subtotal (95% CI)
0.000 [1.20, 0.00)	10000 /0				$ (P = 0.05); ^2$		Heterogeneity: $Tau^2 = 0.1$
					. (Test for overall effect: Z =
						hasone dipropionate	2 Potent steroid: betamet
-1.23 [-1.43, -1.03	100.0 %		-2.77 (1.8)	135	-4.59 (1.39)	550	Jemec 2008 (P)
-1.23 [-1.43, -1.03	100.0 %	•		135		550	Subtotal (95% CI)
						able	Heterogeneity: not applica
						12.09 (P < 0.00001)	Test for overall effect: Z =
Not estimabl				0		hasone valerate 0	3 Potent steroid: betamet Subtotal (95% CI)
						able	Heterogeneity: not applica
						applicable	Test for overall effect: not
						cinonide	4 Very potent steroid: am
-0.97 [-1.33, -0.61	100.0 %		-1.33 (0.89)	67	-2.2 (0.89)	65	Ellis 1988
-0.97 [-1.33, -0.61	100.0 %	•		67		65	Subtotal (95% CI)
							Heterogeneity: not applica
						· · · · · ·	Test for overall effect: Z =
Not estimabl				0		betasol propionate 0	5 Very potent steroid: clo Sb+o+ol (050/, CI)
Not estimadi				U			Subtotal (95% CI) Heterogeneity: not applica
							Test for overall effect: not
							6 Very potent steroid: hal
Not estimabl				0		0	Subtotal (95% CI)
						able	Heterogeneity: not applica
						applicable	Test for overall effect: not
)	on: calcipotriol + BME	7 Vitamin D in combinatio
-1.39 [-1.59, -1.19	51.7 %	-	-2.77 (1.8)	135	-4.75 (1.31)	529	Jemec 2008 (P)
-0.59 [-0.94, -0.24	48.3 %	-	1.93 (1.26)	42	1.3 (0.99)	135	Tyring 2010
	100.0 %			177		664	Subtotal (95% CI)

(Continued . . .)

Study or subgroup	Active treatment	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	(Continued) Std. Mean Difference IV,Random,95% Cl
Heterogeneity: Tau ² = 0.30		· · /		riean(SD)	IV,Rahuom,75% Ci		IV,Nandom,73% CI
Test for overall effect: $Z =$		1 (P - 0.00012)	; 1~ -93%				
8 Other treatment: betame	. ,	nionate plus salio	vlic acid				
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applical	ble						
Test for overall effect: not a							
9 Other treatment: ciclopir	rox olamine shampoo)					
Shuttleworth 1998	28	-2 (1)	9	-1.89 (0.93)	-	100.0 %	-0.11 [-0.86, 0.64]
Subtotal (95% CI)	28		9		•	100.0 %	-0.11 [-0.86, 0.64]
Heterogeneity: not applical	ble						
Test for overall effect: $Z =$	0.29 (P = 0.78)						
10 Other treatment: fluoci	nolone acetonide, pli	us occlusion					
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applical	ble						
Test for overall effect: not a	applicable						
II Other treatment: salicyl	ic acid						
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applical	ble						
Test for overall effect: not a	applicable						
Test for subgroup difference	es: $Chi^2 = 10.67$, df	= 4 (P = 0.03), I ²	=63%				
				-4	-2 0 2	4	

Favours active treatment

Favours placebo

Analysis 18.5. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Active treatment N	Mean(SD)	Placebo N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
25	-2.52 (1.08)	24	-1.25 (1.22)		37.9 %	-1.09 [-1.69, -0.48]
272	1.85 (1.15)	136	2.41 (1.06)		62.1 %	-0.50 [-0.71, -0.29]
297		160		•	100.0 %	-0.72 [-1.28, -0.16]
$hi^2 = 3.25, df = 1$ ((P = 0.011) me dipropionate	$P = 0.07$); $I^2 = 0$	69%				
10	-1.8 (1)	10	-0.9 (1)		4.3 %	-0.86 [-1.79, 0.06]
556	1.23 (1.07)	136	2.41 (1.06)	-	95.7 %	-1.10 [-1.30, -0.91]
566 ² = 0.25, df = 1 (P	= 0.62); l ² =0.	146 0%		•	100.0 %	-1.09 [-1.29, -0.90]
5 (P < 0.00001) ne valerate	2 70 (2 27)			_		
	2./8 (2.2/)		5.97 (2.27)			-1.40 [-1.75, -1.05]
(P < 0.00001) ide 65	-4.6 (1.29)	57 67	-2.8 (1.23)	•	100.0 %	-1.40 [-1.75, -1.05] -1.42 [-1.80, -1.04
65	~ /	67	()	•	100.0 %	-1.42 [-1.80, -1.04]
(P < 0.00001) ol propionate		07			100.0 /0	-1.12 [-1.00, -1.01]
7317 (0.928545)	41	40	2.6 (0.955416)		14.2 %	-1.99 [-2.53, -1.45]
125	2.1 (2.27)	63	5.4 (2.27)	-	26.3 %	-1.45 [-1.78, -1.11]
95	3.2 (1.74)	47	5.49 (1.74)	-	22.7 %	-1.31 [-1.69, -0.93]
188	-3.65 (1.24)	189	-1.7 (1.09)	-	36.7 %	-1.67 [-1.90, -1.43]
440		339		•	100.0 %	-1.57 [-1.81, -1.34]
	N 25 272 297 $i^2 = 3.25, df = 1$ ((P = 0.011)) i^2 dipropionate 10 566 566 566 566 $i^2 = 0.25, df = 1$ (P 5 (P < 0.00001) i^2 dipropionate 65 (P < 0.00001) i^2 dipropionate 7317 (0.928545) 125 95	N Mean(SD) 25 -2.52 (1.08) 272 1.85 (1.15) 297 -1.85 (1.15) i² = 3.25, df = 1 (P = 0.07); l² = 0. (P = 0.011) ae dipropionate 10 -1.8 (1) 556 1.23 (1.07) 566 $e 0.25$, df = 1 (P = 0.62); l² = 0. 5 (P < 0.00001)	N Mean(SD) N 25 -2.52 (1.08) 24 272 1.85 (1.15) 136 297 160 i² = 3.25, df = 1 (P = 0.07); l² = 69% (P = 0.011) i² = 3.25, df = 1 (P = 0.07); l² = 69% 10 i0 -1.8 (1) 10 556 1.23 (1.07) 136 566 146 6 566 9.05, df = 1 (P = 0.62); l² = 0.0% 57 115 2.78 (2.27) 57 115 2.78 (2.27) 57 115 57 57 (P < 0.00001)	N Mean(SD) N Mean(SD) 25 -2.52 (1.08) 24 -1.25 (1.22) 272 1.85 (1.15) 136 2.41 (1.06) 297 1.60 - - $i^2 = 3.25, df = 1 (P = 0.07); l^2 = 69%$ (P = 0.011) 10 -0.9 (1) ie dipropionate 10 -1.8 (1) 10 -0.9 (1) 556 1.23 (1.07) 136 2.41 (1.06) 566 146 - - e 0.25, df = 1 (P = 0.62); l^2 = 0.0% - - - 5 (P < 0.00001)	Active treatment Placebo Mean(SD) N Mean(SD) Mean(SD) 25 -2.52 (1.08) 24 -1.25 (1.22) $-$ 272 1.85 (1.15) 136 2.41 (1.06) $ 297$ 160 $ 297$ 160 $ a^2 = 3.25$, $df = 1$ ($P = 0.07$); $l^2 = 69%$ $ (P = 0.01)$ $ -1.8$ (1) 10 -0.9 (1) $a c d propionate$ $ -1.8$ (1) 10 -0.9 (1) $a c d propionate$ $ a c d propionate$ $ (P < 0.00001)$ $ a c d propionate$ $ -$	Active treatment Placebo Difference Weight N Mean(SD) N Mean(SD) IVRandom,95% CI 25 -2.52 (1.08) 24 -1.25 (1.22) - 37.9 % 272 1.85 (1.15) 136 2.41 (1.06) - 62.1 % 297 160 - 100.0 % 62.1 % 297 160 - 43.3 % 256 1.23 (1.07) 136 2.41 (1.06) 95.7 % 566 146 - 100.0 % 95.7 % 566 146 - 100.0 % 95.7 % 566 146 - 100.0 % 95.7 % 566 146 - 100.0 % 95.7 % 566 146 - 100.0 % 95.7 % 57 597 (2.27) I000.0 % 100.0 % 65 67 - 1000.0 % 65 67 - 100.0 % 65 67 - 100.0 %

(Continued . . .)

Study or subgroup Act	ive treatment		Placebo		Std. Mean Difference	Weight	(Continue) Std Mear Difference
, , ,	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% Cl	-	IV,Random,95% C
6 Very potent steroid: halcinonide							
Lepaw 1978	27	-2.3 (0.95)	27	-1.3 (0.82)	H	100.0 %	-1.11 [-1.69, -0.53]
Subtotal (95% CI)	27		27		•	100.0 %	-1.11 [-1.69, -0.53]
Heterogeneity: not applicable							
Test for overall effect: $Z = 3.78$ (P	= 0.00016)						
7 Vitamin D in combination: calcip	otriol + BMD				_		
Jemec 2008 (P)	541	1.09 (1.02)	136	2.41 (1.06)	•	52.5 %	-1.28 [-1.48, -1.08
Tyring 2010	135	1.05 (1.15)	42	1.79 (1.28)	-	47.5 %	-0.62 [-0.98, -0.27
Subtotal (95% CI)	676		178		•	100.0 %	-0.97 [-1.61, -0.32]
Heterogeneity: Tau ² = 0.20; Chi ² = Test for overall effect: Z = 2.95 (P 8 Other treatment: betamethason	= 0.0032)						
Elie 1983	10	-2.45 (1)	10	-0.9 (1)		100.0 %	-1.48 [-2.50, -0.47
Subtotal (95% CI)	10		10		-	100.0 %	-1.48 [-2.50, -0.47
Heterogeneity: not applicable Test for overall effect: Z = 2.86 (P 9 Other treatment: ciclopirox olarr	,						
Shuttleworth 1998	28	4.25 (2.49)	9	4.44 (2.65)	-	100.0 %	-0.07 [-0.82, 0.68
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 0.19 (P	28 = 0.85)		9		•	1 00.0 %	-0.07 [-0.82, 0.68
10 Other treatment: fluocinolone a	acetonide, plus	occlusion					
Pauporte 2004	42	-4.38 (0.99)	42	-2.76 (1.57)	-	100.0 %	-1.22 [-1.69, -0.76
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 5.12 (P	42 < 0.00001)		42		•	1 00.0 %	-1.22 [-1.69, -0.76]
I I Other treatment: salicylic acid							
Elie 1983	10	-1.8 (1)	10	-0.9 (1)		100.0 %	-0.86 [-1.79, 0.06
Subtotal (95% CI) Heterogeneity: not applicable Test for overall effect: Z = 1.82 (P Test for subgroup differences: Chi ²	10 = 0.068)		10		-	100.0 %	-0.86 [-1.79, 0.06

Favours active treatment Favours placebo

Analysis 18.6. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: 6 Total withdrawals

Study or subgroup	Active treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
I Vitamin D: calcipotriol					
Green 1994	1/25	2/24		23.2 %	-0.04 [-0.18, 0.09]
Jemec 2008 (P)	57/272	30/136	-	58.5 %	-0.01 [-0.10, 0.07]
Kiss 1996	3/30	3/30	_	18.3 %	0.0 [-0.15, 0.15]
Subtotal (95% CI)	327	190	+	100.0 %	-0.02 [-0.08, 0.05]
Total events: 61 (Active treatm Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: Z = 0.50 2 Potent steroid: betamethasor	$^{2} = 0.23$, df = 2 (P = 0.8 0 (P = 0.62)	9); I ² =0.0%			
Jemec 2008 (P)	47/556	30/136	H	100.0 %	-0.14 [-0.21, -0.06]
Subtotal (95% CI)	556	136	•	100.0 %	-0.14 [-0.21, -0.06]
Heterogeneity: not applicable Test for overall effect: Z = 3.63 3 Potent steroid: betamethasor Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not appli	ne valerate 0 nt), 0 (Placebo)	0			Not estimable
4 Very potent steroid: amcinon					
Ellis 1988	13/83	13/82		100.0 %	0.00 [-0.11, 0.11]
Subtotal (95% CI) Total events: 13 (Active treatme		82	-	100.0 %	0.00 [-0.11, 0.11]
Heterogeneity: not applicable Test for overall effect: Z = 0.03 5 Very potent steroid: clobetas	· /				
Test for overall effect: $Z = 0.03$	· /	2/40		14.9 %	0.05 [-0.07, 0.16]
Test for overall effect: Z = 0.03 5 Very potent steroid: clobetas	ol propionate	2/40 0/63		14.9 % 24.3 %	0.05 [-0.07, 0.16] 0.0 [-0.02, 0.02]
Test for overall effect: Z = 0.03 5 Very potent steroid: clobetas Cook-Bolden 2010	ol propionate 4/41		+-		2 3
Test for overall effect: Z = 0.03 5 Very potent steroid: clobetas Cook-Bolden 2010 Franz 2000	ol propionate 4/41 0/125	0/63	 • •	24.3 %	0.0 [-0.02, 0.02]

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Study or subgroup	Active treatment	Placebo	Difference M- H,Random,95%	Weight	Difference M H,Random,
	n/N	n/N	Ćl		C
Subtotal (95% CI)	556	450	•	100.0 %	-0.03 [-0.10, 0.04]
Total events: 26 (Active treat	, , ,				
Heterogeneity: $Tau^2 = 0.00;$		00001); l ² =89%			
Test for overall effect: $Z = 0.7$. ,				
6 Very potent steroid: halcing		2 /20			
Lepaw 1978	2/29	2/29		100.0 %	0.0 [-0.13, 0.13
Subtotal (95% CI)	29	29	-	100.0 %	0.0 [-0.13, 0.13]
Total events: 2 (Active treatm	nent), 2 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
7 Vitamin D in combination:	calcipotriol + BMD				
Jemec 2008 (P)	61/541	30/136		75.9 %	-0.11 [-0.18, -0.03]
Tyring 2010	19/135	8/42		24.1 %	-0.05 [-0.18, 0.08
Subtotal (95% CI)	676	178	•	100.0 %	-0.09 [-0.16, -0.03]
Total events: 80 (Active treat	ment), 38 (Placebo)				
Heterogeneity: Tau ² = 0.0; C	, , ,	5): $ ^2 = 0.0\%$			
Test for overall effect: $Z = 2.1$,	· //			
3 Other treatment: betameth	, ,	plus salicylic acid			
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Active treatm	nent), 0 (Placebo)				
Heterogeneity: not applicable	2				
Test for overall effect: not ap	olicable				
Other treatment: ciclopiro>	olamine shampoo				
Shuttleworth 1998	1/29	2/11		100.0 %	-0.15 [-0.38, 0.09
Subtotal (95% CI)	29	11		100.0 %	-0.15 [-0.38, 0.09
Total events: (Active treatm	nent), 2 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 1.2$	22 (P = 0.22)				
10 Other treatment: fluocinc	lone acetonide, plus occlus	ion			
Pauporte 2004	1/43	3/46		100.0 %	-0.04 [-0.13, 0.04
Subtotal (95% CI)	43	46	•	100.0 %	-0.04 [-0.13, 0.04
Total events: 1 (Active treatm					
-leterogeneity: not applicable	, , ,				
Test for overall effect: $Z = 0.9$					
	· · ·				
I Other treatment: salicylic		0			Nat antim-11
Subtotal (95% CI)	0	0			Not estimable
Tatal suggests O (A 2) and a	, , ,				
Total events: 0 (Active treatm	2				
Heterogeneity: not applicable					
Heterogeneity: not applicable Test for overall effect: not ap	olicable	0 17) 12 -270/			
Heterogeneity: not applicable	olicable	0.17), 12 =32%			

Analysis 18.7. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Active treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95 Cl
I Vitamin D: calcipotriol					
Green 1994	1/25	0/24	-	11.7 %	0.04 [-0.07, 0.15]
Jemec 2008 (P)	20/272	7/136	•	55.3 %	0.02 [-0.03, 0.07]
Kiss 1996	0/30	0/30	+	32.9 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	327	190	•	100.0 %	0.02 [-0.02, 0.05]
Total events: 21 (Active treat	ment), 7 (Placebo)				
Heterogeneity: Tau ² = 0.0; C	² hi ² = 0.56, df = 2 (P = 0.7	6); l ² =0.0%			
Test for overall effect: $Z = 0.9$	92 (P = 0.36)				
2 Potent steroid: betamethas	one dipropionate				
Elie 1983	0/10	0/10		4.6 %	0.0 [-0.17, 0.17]
Jemec 2008 (P)	6/556	7/136	-	95.4 %	-0.04 [-0.08, 0.00]
Subtotal (95% CI)	566	146	•	100.0 %	-0.04 [-0.08, 0.00]
Total events: 6 (Active treatm	nent), 7 (Placebo)				
Heterogeneity: Tau ² = 0.0; C	$hi^2 = 0.20, df = 1 (P = 0.6)$	5); l ² =0.0%			
Test for overall effect: $Z = 2.0$	04 (P = 0.041)				
3 Potent steroid: betamethas	one valerate				
Franz 1999	0/115	0/57		100.0 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	115	57	•	100.0 %	0.0 [-0.03, 0.03]
Total events: 0 (Active treatm	nent), 0 (Placebo)				
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.0$	0 (P = 1.0)				
4 Very potent steroid: amcine	onide				
Ellis 1988	1/83	0/82	•	100.0 %	0.01 [-0.02, 0.04]
Subtotal (95% CI)	83	82	•	100.0 %	0.01 [-0.02, 0.04]
Total events: I (Active treatm	nent), 0 (Placebo)				
Heterogeneity: not applicable	2				
Test for overall effect: $Z = 0.7$	72 (P = 0.47)				
			-1 -0.5 0 0.5 1		
			-1 -0.5 0 0.5 1 tive treatment Favours placeb	0	
		i avoui s de	ave a cauncile in avoid s placeb		(Continued)

(Continued . . .)

(... Continued) Risk Risk Difference Difference Study or subgroup Active treatment Placebo Weight M-M-H,Random,95% H,Random,95% n/N n/N ĆΙ Ć 5 Very potent steroid: clobetasol propionate -0.03 [-0.09, 0.04] Cook-Bolden 2010 0/41 1/40 2.8 % Franz 2000 0/125 20.9 % 0.0 [-0.02, 0.02] 0/63 Jarratt 2004 0/95 0/47 11.9 % 0.0 [-0.03, 0.03] Olsen 1991 0/189 57.9 % 1/189 -0.01 [-0.02, 0.01] Poulin 2010 2/106 4/||| 6.5 % -0.02 [-0.06, 0.03] Subtotal (95% CI) 556 450 100.0 % 0.00 [-0.02, 0.01] Total events: 2 (Active treatment), 6 (Placebo) Heterogeneity: Tau² = 0.0; Chi² = 1.22, df = 4 (P = 0.88); $I^2 = 0.0\%$ Test for overall effect: Z = 0.87 (P = 0.39) 6 Very potent steroid: halcinonide 0.0 [-0.06, 0.06] Lepaw 1978 0/29 0/29 100.0 % Subtotal (95% CI) 29 29 100.0 % 0.0 [-0.06, 0.06] Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0) 7 Vitamin D in combination: calcipotriol + BMD Jemec 2008 (P) 8/541 7/136 100.0 % -0.04 [-0.08, 0.00] Subtotal (95% CI) 100.0 % -0.04 [-0.08, 0.00] 541 136 Total events: 8 (Active treatment), 7 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 1.87 (P = 0.062) 8 Other treatment: betamethasone-17,21-dipropionate plus salicylic acid Elie 1983 0/10 0/10 100.0 % 0.0 [-0.17, 0.17] Subtotal (95% CI) 100.0 % 0.0 [-0.17, 0.17] 10 10 Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0)9 Other treatment: ciclopirox olamine shampoo Shuttleworth 1998 2/11 100.0 % -0.18 [-0.42, 0.05] 0/29 Subtotal (95% CI) 29 11 100.0 % -0.18 [-0.42, 0.05] Total events: 0 (Active treatment), 2 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 1.52 (P = 0.13) 10 Other treatment: fluocinolone acetonide, plus occlusion Subtotal (95% CI) 0 0 Not estimable Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: not applicable - [-0.5 0 0.5 Favours active treatment Favours placebo

(Continued ...)

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Active treatment	Placebo n/N	Risk Difference M. H,Random,95% Cl	Weight	(Continued) Risk Difference H,Random,95%
II Other treatment: salicylic		11/14			CI
Elie 1983	0/10	0/10	-	100.0 %	0.0 [-0.17, 0.17]
Subtotal (95% CI)	10	10	+	100.0 %	0.0 [-0.17, 0.17]
Total events: 0 (Active treat	ment), 0 (Placebo)				
Heterogeneity: not applicab	le				
Test for overall effect: $Z = 0$	0.0 (P = 1.0)				
Test for subgroup difference	es: $Chi^2 = 10.57$, $df = 9$ (P =	0.3 I), I ² = I 5%			
			-1 -0.5 0 0.5 1		
		Favours a	active treatment Favours placebo)	

Analysis 18.8. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Active treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
I Vitamin D: calcipotriol					
Green 1994	0/25	2/24		18.8 %	-0.08 [-0.21, 0.05]
Jemec 2008 (P)	19/272	16/136	=	81.2 %	-0.05 [-0.11, 0.01]
Subtotal (95% CI)	297	160	•	100.0 %	-0.05 [-0.11, 0.00]
Total events: 19 (Active trea	tment), 18 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$; ($Chi^2 = 0.24, df = 1 (P = 0.6)$	52); I ² =0.0%			
Test for overall effect: $Z = I$.91 (P = 0.056)				
2 Potent steroid: betametha	sone dipropionate				
Jemec 2008 (P)	9/556	16/136		100.0 %	-0.10 [-0.16, -0.05]
Subtotal (95% CI)	556	136	•	100.0 %	-0.10 [-0.16, -0.05]
		F	-0.5 -0.25 0 0.25 0.5		
		Favou	rs active treatment Favours placebo		

(Continued . . .)

Ci la la	A		Risk	NA / 1 /	Risk
Study or subgroup	Active treatment	Placebo	Difference M-	Weight	Difference M
	n/N	n/N	H,Random,95% Cl		H,Random, C
Total events: 9 (Active treatr	ment), 16 (Placebo)				
Heterogeneity: not applicabl	e				
Test for overall effect: $Z = 3$.	.61 (P = 0.00031)				
3 Potent steroid: betametha	sone valerate				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Active treatr	, , ,				
Heterogeneity: not applicabl					
Test for overall effect: not ap					
4 Very potent steroid: amcin		1 (00)			
Ellis 1988	0/83	1/82		100.0 %	-0.01 [-0.05, 0.02
Subtotal (95% CI)	83	82	+	100.0 %	-0.01 [-0.05, 0.02
Total events: 0 (Active treatr	, , ,				
Heterogeneity: not applicabl					
Test for overall effect: $Z = 0$.	, ,				
5 Very potent steroid: clobe Cook-Bolden 2010	tasol propionate 0/41	0/40	-	17.3 %	
COOK-BOIDEN 2010	0/41	0/40	T	17.3 /0	0.0 [-0.05, 0.05
Franz 2000	0/125	0/63	•	22.2 %	0.0 [-0.02, 0.02
Jarratt 2004	0/95	0/47	+	20.6 %	0.0 [-0.03, 0.03
Olsen 1991	2/189	17/189	-	18.1 %	-0.08 [-0.12, -0.04
Poulin 2010	1/106	0/111	-	21.9 %	0.01 [-0.02, 0.03
Subtotal (95% CI)	556	450	•	100.0 %	-0.01 [-0.05, 0.02
Total events: 3 (Active treatr	ment), I7 (Placebo)				
Heterogeneity: $Tau^2 = 0.00;$	$Chi^2 = 23.90, df = 4 (P = 0)$	0.00008); l ² =83%			
Test for overall effect: $Z = 0$.	, ,				
6 Very potent steroid: halcin					
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Active treatr	, , , ,				
Heterogeneity: not applicabl Test for overall effect: not ap					
7 Vitamin D in combination:					
Jemec 2008 (P)	2/541	16/136	-	100.0 %	-0.11 [-0.17, -0.06
		126	•		-
Subtotal (95% CI)	541	136	•	100.0 %	-0.11 [-0.17, -0.06]
Total events: 2 (Active treatr Heterogeneity: not applicabl	, , ,				
Test for overall effect: $Z = 4$.					
8 Other treatment: betamet	. ,	plus salicylic acid			
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Active treatr					
Heterogeneity: not applicabl	e				
Test for overall effect: not ap	plicable				
			0.5 -0.25 0 0.25 0.5		
		-(0.0 -0.25 0 0.25 0.5		

Study or subgroup	Active treatment Placebo		Risk Difference H.Random,95%	Weight	(Continued) Risk Difference M- H.Random,959	
	n/N	n/N	Ċl		Ċl	
9 Other treatment: ciclopirox of	1					
Shuttleworth 1998	0/29	1/11		100.0 %	-0.09 [-0.28, 0.10]	
Subtotal (95% CI)	29	11		100.0 %	-0.09 [-0.28, 0.10]	
Total events: 0 (Active treatme	ent), I (Placebo)					
Heterogeneity: not applicable						
Test for overall effect: $Z = 0.92$	2 (P = 0.36)					
10 Other treatment: fluocinolo	one acetonide, plus occlus	ion				
Subtotal (95% CI)	0	0			Not estimable	
Subtotal (95% CI) Total events: 0 (Active treatme	-	0			Not estimable	
, , ,	-	0			Not estimable	
Total events: 0 (Active treatme Heterogeneity: not applicable	ent), 0 (Placebo)	0			Not estimable	
Total events: 0 (Active treatme Heterogeneity: not applicable Test for overall effect: not appl	ent), 0 (Placebo) iicable	0			Not estimable	
Total events: 0 (Active treatme Heterogeneity: not applicable Test for overall effect: not appl 11 Other treatment: salicylic ac	ent), 0 (Placebo) iicable	0			Not estimable Not estimable	
Total events: 0 (Active treatme Heterogeneity: not applicable Test for overall effect: not appl 11 Other treatment: salicylic ac Subtotal (95% CI)	ent), 0 (Placebo) icable cid 0	-				
Total events: 0 (Active treatme Heterogeneity: not applicable Test for overall effect: not appli 11 Other treatment: salicylic ac Subtotal (95% CI) Total events: 0 (Active treatme	ent), 0 (Placebo) icable cid 0	-				
Total events: 0 (Active treatme Heterogeneity: not applicable Test for overall effect: not appli 11 Other treatment: salicylic ac Subtotal (95% CI) Total events: 0 (Active treatme Heterogeneity: not applicable	ent), 0 (Placebo) iicable cid o ent), 0 (Placebo)	-				
Total events: 0 (Active treatme Heterogeneity: not applicable Test for overall effect: not appli 11 Other treatment: salicylic ac Subtotal (95% CI) Total events: 0 (Active treatme	ent), 0 (Placebo) iicable cid ent), 0 (Placebo) iicable	0				

Favours active treatment Favours placebo

Analysis 18.9. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: 9 Adverse events (local)

Study or subgroup	Active treatment	Placebo	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random, Cl
I Vitamin D: calcipotriol					
Green 1994	5/25	7/24		3.7 %	-0.09 [-0.33, 0.15]
Jemec 2008 (P)	35/266	18/135	-	42.7 %	0.00 [-0.07, 0.07]
Kiss 1996	0/30	0/30	+	53.6 %	0.0 [-0.06, 0.06]
Subtotal (95% CI)	321	189	•	100.0 %	0.00 [-0.05, 0.04]
Total events: 40 (Active treatme	nt), 25 (Placebo)				
Heterogeneity: $Tau^2 = 0.0$; Chi ²	= 0.62, df = 2 (P = 0.72	3); I ² =0.0%			
Test for overall effect: $Z = 0.18$	(P = 0.86)				
2 Potent steroid: betamethason	e dipropionate				
Elie 1983	0/10	0/10		10.7 %	0.0 [-0.17, 0.17]
Jemec 2008 (P)	29/548	18/135	-	89.3 %	-0.08 [-0.14, -0.02]
Subtotal (95% CI)	558	145	•	100.0 %	-0.07 [-0.13, -0.01]
Total events: 29 (Active treatme	nt), 18 (Placebo)				
	-0.7(-1)(-0.2)	0), 12 -0.0%			
Heterogeneity: Tau ² = 0.0; Chi ²	-0.76, dt -1 (P -0.36	b), 1 =0.076			
Heterogeneity: Tau ² = 0.0; Chi ² Test for overall effect: $Z = 2.47$		5), 1 -0.076			
Test for overall effect: $Z = 2.47$	(P = 0.014)	5), 1 -0.078			
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone	(P = 0.014)	0			Not estimable
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI)	(P = 0.014) e valerate 0	,			Not estimable
0,	(P = 0.014) e valerate 0	,			Not estimable
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable	(P = 0.014) e valerate t), 0 (Placebo)	,			Not estimable
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic	(P = 0.014) e valerate t), 0 (Placebo) able	,			Not estimable
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinoni	(P = 0.014) e valerate t), 0 (Placebo) able	,			
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinoni Subtotal (95% CI)	(P = 0.014) e valerate t), 0 (Placebo) able de 0	0			
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinonie Subtotal (95% CI) Total events: 0 (Active treatmen	(P = 0.014) e valerate t), 0 (Placebo) able de 0	0			
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinonic Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable	(P = 0.014) e valerate t), 0 (Placebo) able de t), 0 (Placebo)	0			
Test for overall effect: Z = 2.47 8 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinonic Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applicable	(P = 0.014) e valerate 0 t), 0 (Placebo) able de 0 t), 0 (Placebo) able	0			
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinonic Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic	(P = 0.014) e valerate 0 t), 0 (Placebo) able de 0 t), 0 (Placebo) able	0		19.6 %	Not estimable
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinonic Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 5 Very potent steroid: clobetasc	(P = 0.014) e valerate 0 t), 0 (Placebo) able de 0 t), 0 (Placebo) able	0 0		19.6 % 7.1 %	Not estimable 0.02 [-0.06, 0.11]
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinonic Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 5 Very potent steroid: clobetasc Cook-Bolden 2010	(P = 0.014) e valerate 0 t), 0 (Placebo) able de 0 t), 0 (Placebo) able ol propionate 2/41	0 0 1/40			Not estimable Not estimable 0.02 [-0.06, 0.11] -0.07 [-0.21, 0.06] 0.02 [-0.05, 0.08]
Test for overall effect: Z = 2.47 3 Potent steroid: betamethasone Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 4 Very potent steroid: amcinoni Subtotal (95% CI) Total events: 0 (Active treatmen Heterogeneity: not applicable Test for overall effect: not applic 5 Very potent steroid: clobetasc Cook-Bolden 2010 Jarratt 2004	(P = 0.014) e valerate 0 t), 0 (Placebo) able de 0 t), 0 (Placebo) able ol propionate 2/41 13/94	0 0 1/40 10/47		7.1 %	Not estimable 0.02 [-0.06, 0.1 -0.07 [-0.21, 0.06

(Continued \dots)

(... Continued) Risk Risk Difference Difference Study or subgroup Active treatment Placebo Weight M-M-H,Random,95% H,Random,95% n/N n/N Ć Ć Total events: 41 (Active treatment), 35 (Placebo) Heterogeneity: Tau² = 0.0; Chi² = 1.87, df = 3 (P = 0.60); $I^2 = 0.0\%$ Test for overall effect: Z = 0.12 (P = 0.90) 6 Very potent steroid: halcinonide Lepaw 1978 0/29 1/29 100.0 % -0.03 [-0.12, 0.06] Subtotal (95% CI) 29 29 100.0 % -0.03 [-0.12, 0.06] Total events: 0 (Active treatment), | (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.75 (P = 0.45) 7 Vitamin D in combination: calcipotriol + BMD Jemec 2008 (P) 25/530 18/135 62.1 % -0.09 [-0.15, -0.03] Tyring 2010 9/128 3/38 37.9 % -0.01 [-0.11, 0.09] Subtotal (95% CI) 658 173 100.0 % -0.06 [-0.13, 0.02] Total events: 34 (Active treatment), 21 (Placebo) Heterogeneity: Tau² = 0.00; Chi² = 1.82, df = 1 (P = 0.18); $I^2 = 45\%$ Test for overall effect: Z = 1.49 (P = 0.14)8 Other treatment: betamethasone-17,21-dipropionate plus salicylic acid 0.0 [-0.17, 0.17] Elie 1983 0/10 0/10 100.0 % Subtotal (95% CI) 10 10 0.0 [-0.17, 0.17] 100.0 % Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.0 (P = 1.0) 9 Other treatment: ciclopirox olamine shampoo Shuttleworth 1998 1/29 1/11 100.0 % -0.06 [-0.24, 0.13] Subtotal (95% CI) -0.06 [-0.24, 0.13] 29 11 100.0 % Total events: | (Active treatment), | (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.61 (P = 0.54) 10 Other treatment: fluocinolone acetonide, plus occlusion Pauporte 2004 1/43 0/46 100.0 % 0.02 [-0.04, 0.08] Subtotal (95% CI) 0.02 [-0.04, 0.08] 43 100.0 % 46 Total events: I (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: Z = 0.75 (P = 0.46) II Other treatment: salicylic acid Elie 1983 0/10 0/10 100.0 % 0.0 [-0.17, 0.17] Subtotal (95% CI) 10 10 100.0 % 0.0 [-0.17, 0.17] Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable -05 -0.25 0 025 05 Favours active treatment Favours placebo

(Continued . . .)

Topical treatments for chronic plaque psoriasis (Review)

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						(Continued)
Study or subgroup	Active treatment	Placebo		Risk erence M-	Weight	Risk Difference M-
	n/N	n/N	H,Ra	ndom,95% Cl		H,Random,95% Cl
Test for overall effect: $Z = 0$).0 (P = 1.0)					
Test for subgroup difference	es: $Chi^2 = 8.05$, $df = 8$ (P = 0	0.43), I ² =I%				
			<u> </u>			
			-0.5 -0.25	0 0.25 0.5	5	
		Favours a	active treatment	Favours place	bo	

Analysis 18.10. Comparison 18 Scalp psoriasis: placebo-controlled trials, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 18 Scalp psoriasis: placebo-controlled trials

Outcome: 10 Adverse events (systemic)

Study or subgroup	Active treatment	Placebo	Risk Difference M- H,Random,95%	Weight	Risk Difference M-
	n/N	n/N	H,Kandom,95% Cl		H,Random,95% Cl
I Vitamin D: calcipotriol					
Jemec 2008 (P)	0/272	0/136	-	100.0 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	272	136	•	100.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Active treatm	ent), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$) (P = 1.0)				
2 Potent steroid: betamethas	one dipropionate				
Jemec 2008 (P)	0/556	0/136	-	100.0 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	556	136	•	100.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Active treatm	ient), 0 (Placebo)				
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
3 Potent steroid: betamethas	one valerate				
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Active treatm	ient), 0 (Placebo)				
Heterogeneity: not applicable					
			-0.2 -0.1 0 0.1 0.2		
		Favou	rs active treatment Favours placeb	0	

(Continued . . .)

Topical treatments for chronic plaque psoriasis (Review)

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(... Continued) Risk Risk Difference Difference Study or subgroup Active treatment Placebo Weight M-M-H,Random,95% H,Random,95% n/N n/N ĆΙ Ć Test for overall effect: not applicable 4 Very potent steroid: amcinonide Subtotal (95% CI) 0 0 Not estimable Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: not applicable 5 Very potent steroid: clobetasol propionate Olsen 1991 4/83 4/85 13.0 % 0.00 [-0.06, 0.07] Poulin 2010 0/106 1/111 87.0 % -0.01 [-0.03, 0.02] Subtotal (95% CI) 189 196 100.0 % -0.01 [-0.03, 0.02] Total events: 4 (Active treatment), 5 (Placebo) Heterogeneity: Tau² = 0.0; Chi² = 0.15, df = 1 (P = 0.70); l² = 0.0% Test for overall effect: Z = 0.65 (P = 0.52) 6 Very potent steroid: halcinonide Subtotal (95% CI) 0 Not estimable 0 Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: not applicable 7 Vitamin D in combination: calcipotriol + BMD Jemec 2008 (P) 0/541 0/136 92.6 % 0.0 [-0.01, 0.01] Tyring 2010 0/128 0/38 74% 0.0 [-0.04, 0.04] Subtotal (95% CI) 669 174 100.0 % 0.0 [-0.01, 0.01] Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: Tau² = 0.0; Chi² = 0.0, df = 1 (P = 1.00); l² = 0.0% Test for overall effect: Z = 0.0 (P = 1.0) 8 Other treatment: betamethasone-17,21-dipropionate plus salicylic acid Subtotal (95% CI) 0 Not estimable 0 Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: not applicable 9 Other treatment: ciclopirox olamine shampoo Subtotal (95% CI) 0 0 Not estimable Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: not applicable 10 Other treatment: fluocinolone acetonide, plus occlusion Not estimable Subtotal (95% CI) 0 0 Total events: 0 (Active treatment), 0 (Placebo) Heterogeneity: not applicable Test for overall effect: not applicable II Other treatment: salicylic acid -0.2 -0.1 0 0.1 0.2 Favours active treatment

Favours placebo

(Continued ...)

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Active treatment	Placebo		te 1-	Weight	(Continued) Risk Difference M-
	n/N	n/N	H,Random,95% Cl			H,Random,95% Cl
Subtotal (95% CI)	0	0				Not estimable
Total events: 0 (Active treatm	nent), 0 (Placebo)					
Heterogeneity: not applicable	e					
Test for overall effect: not ap	plicable					
Test for subgroup differences	s: $Chi^2 = 0.39$, $df = 3$ (P = 0	0.94), l ² =0.0%				
			-0.2 -0.1 0	0.1 0.2		
		Favours a	ctive treatment F	avours placebo		

Analysis 19.1. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 1 IAGI.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: I IAGI

Study or subgroup	Vitamin D alone or in combination N	Mean(SD)	Other active treatment N	Mean(SD)	Std. Mean Difference IV,Random,95% Cl	Weight	Std. Mean Difference IV,Random,95% CI
l Vitamin D vs. corticoster	roid (potent): calcipo	otriol vs. BMD					
Jemec 2008 (H)	272	1.85 (1.15)	556	1.23 (1.07)		49.5 %	0.56 [0.42, 0.71]
Van de Kerkhof 2009	286	1.73 (1.1)	562	1.3 (1.07)	-	50.5 %	0.40 [0.25, 0.54]
Subtotal (95% CI)	558		1118		•	100.0 %	0.48 [0.32, 0.64]
Heterogeneity: $Tau^2 = 0.0$	I; Chi ² = 2.53, df =	$ (P = 0.); ^2$	=60%				
Test for overall effect: Z =	5.75 (P < 0.00001)						
2 Vitamin D vs. corticoster	roid (potent): calcipe	otriol vs. BMV					
Duweb 2000	24	-2.63 (0.88)	18	-2.83 (0.92)		8.2 %	0.22 [-0.39, 0.83]
Klaber 1994	236	-2.51 (1.14)	232	-2.93 (1.02)	-	91.8 %	0.39 [0.20, 0.57]
Subtotal (95% CI)	260		250		•	100.0 %	0.37 [0.20, 0.55]
Heterogeneity: $Tau^2 = 0.0;$	$Chi^2 = 0.27, df = 1$	$(P = 0.6 I); I^2 =$	0.0%				
Test for overall effect: Z =	4.18 (P = 0.000029)					
3 Vitamin D vs. corticoster	roid (very potent): c	alcipotriol vs. clc	betasol prop	ionate			
				-2	-I 0 I	2	
			Favours vita	amin D alone or in c	combination Favours ot	her active treatment	
							(Continued

(Continued ...)

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Vitamin D alone or in combination		Other active treatment		Std. Mean Difference	Weight	(Continued Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% CI
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicab	ble						
Test for overall effect: not a	pplicable						
4 Vitamin D + corticostero	id vs. corticosteroio	d: calcipotriol + E	BMD vs. BMI	0			
Buckley 2008	108	0.79 (0.91)	110	1.05 (1.04)		8.9 %	-0.26 [-0.53, 0.00]
Jemec 2008 (H)	541	1.09 (1.02)	556	1.23 (1.07)	-	45.0 %	-0.13 [-0.25, -0.02]
Van de Kerkhof 2009	567	1.08 (1)	562	1.3 (1.07)	=	46.1 %	-0.21 [-0.33, -0.10]
Subtotal (95% CI)	1216		1228		•	100.0 %	-0.18 [-0.26, -0.10]
Heterogeneity: $Tau^2 = 0.0;$	Chi ² = 1.27, df = 2	$(P = 0.53); I^2 =$	0.0%				
Test for overall effect: $Z = 4$	4.48 (P < 0.00001)						
5 Vitamin D vs. vitamin D +	+ corticosteroid: cal	cipotriol vs. calci	potriol + B№	1D			
Jemec 2008 (H)	272	1.85 (1.15)	541	1.09 (1.02)	-	26.4 %	0.71 [0.56, 0.86]
Kragballe 2009	105	2.2 (1.26)	207	1.17 (1.05)	-	21.2 %	0.91 [0.67, 1.16]
Luger 2008	265	1.68 (1.12)	338	1.3 (1.01)	-	25.8 %	0.36 [0.20, 0.52]
Van de Kerkhof 2009	286	1.73 (1.1)	567	1.08 (1)	-	26.6 %	0.63 [0.48, 0.77]
Subtotal (95% CI)	928		1653		•	100.0 %	0.64 [0.44, 0.84]
Heterogeneity: Tau ² = 0.03	; Chi ² = 16.99, df =	= 3 (P = 0.00071); I ² =82%				
Test for overall effect: $Z = e$	6.23 (P < 0.00001)						
6 Vitamin D vs. other treatr	ments: calcipotriol v	rs. coal tar polyth	erapy				
Barrett 2005	161	-3.48 (.)	170	-3.49 (1.17)	-	49.6 %	0.01 [-0.21, 0.22]
Klaber 2000b	208	-2.77 (1.28)	209	-2.07 (1.56)	-	50.4 %	-0.49 [-0.68, -0.29
Subtotal (95% CI)	369		379			100.0 %	-0.24 [-0.73, 0.25]
Heterogeneity: $Tau^2 = 0.11$; Chi ² = 11.30, df =	= I (P = 0.00078); I ² =91%				
Test for overall effect: $Z = 0$	0.97 (P = 0.33)						
Test for subgroup difference	es: Chi ² = 106.49, c	f = 4 (P = 0.00)	, l ² =96%				
				Î		i.	

Analysis 19.2. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 2 TSS.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: 2 TSS

Std Mear Difference IV,Random,95% C	Weight	Std. Mean Difference IV,Random,95% Cl	Mean(SD)	Other active treatment N	Mean(SD)	Vitamin D alone or in combination N	Study or subgroup
					triol vs. BMD	oid (potent): calcipot	I Vitamin D vs. corticoster
0.54 [0.40, 0.69	49.5 %	•	2.2 (2.1)	556	3.4 (2.4)	272	Jemec 2008 (H)
0.36 [0.22, 0.51	50.5 %		2.21 (2)	562	2.95 (2.1)	286	Van de Kerkhof 2009
0.45 [0.28, 0.63]	100.0 %	•		1118		558	Subtotal (95% CI)
				=66%	(P = 0.08); I ²	; Chi ² = 2.97, df = 1	Heterogeneity: Tau ² = 0.01
						5.01 (P < 0.00001)	Test for overall effect: Z =
					triol vs. BMV	oid (potent): calcipot	2 Vitamin D vs. corticoster
0.24 [-0.38, 0.85	8.4 %		1.49 (2.53)	18	2.1 (2.53)	24	Duweb 2000
0.07 [-0.11, 0.26	91.6 %	-	2.93 (4.35)	225	3.29 (5.3)	220	Klaber 1994
0.09 [-0.09, 0.27]	100.0 %	•		243		244	Subtotal (95% CI)
			onate			0.97 (P = 0.33)	Heterogeneity: Tau ² = 0.0; Test for overall effect: Z = 3 Vitamin D vs. corticoster
0.37 [0.05, 0.69	100.0 %		1.76 (1.57)	76	2.36 (1.64)	75	Reygagne 2005
0.37 [0.05, 0.69]	100.0 %	•		76		2.27 (P = 0.024)	Subtotal (95% CI) Heterogeneity: not applicat Test for overall effect: Z =
	8.9 %	-		BMD vs. BMD 110		oid vs. corticosteroid: 108	4 Vitamin D + corticostero
-0.21 [-0.48, 0.05			-4.77 (2.54)		-5.31 (2.47)		Buckley 2008
-0.14 [-0.26, -0.02	45.0 %	-	2.2 (2.1)	556	1.9 (2.1)	541	Jemec 2008 (H)
-0.24 [-0.36, -0.12	46.1 %	•	2.21 (2)	562	1.74 (1.9)	567	Van de Kerkhof 2009
-0.19 [-0.27, -0.11]	100.0 %	•		1228		1216	Subtotal (95% CI)
				0.0%	$(P = 0.5); ^2 =$	Chi ² = 1.36, df = 2	Heterogeneity: $Tau^2 = 0.0$;
						4.79 (P < 0.00001)	Test for overall effect: $Z = $
							5 Vitamin D vs. vitamin D -
0.68 [0.53, 0.83	38.4 %	-	1.9 (2.1)	541	3.4 (2.4)	272	Jemec 2008 (H)
	22.2 %	-	2.3 (2.1)	207	4.4 (2.7)	105	Kragballe 2009
0.90 [0.66, 1.15							
0.90 [0.66, 1.15	39.4 %	•	1.74 (1.9)	567	2.95 (2.1)	286	Van de Kerkhof 2009

Favours vitamin D alone or in combination Favours other active treatment

(Continued ...)

Study or subgroup	Vitamin D alone or in combination		Other active treatment		Di	Std. Mean fference	Weight	(Continued) Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Ranc	lom,95% Cl		IV,Random,95% CI
Heterogeneity: Tau ² = 0.01;	Chi ² = 3.97, df =	2 (P = 0.14); I ²	=50%					
Test for overall effect: $Z = 9$.78 (P < 0.00001)							
6 Vitamin D vs. other treatm	nents: calcipotriol v	s. coal tar polyt	herapy					
Barrett 2005	207	-2.8 (2.3)	211	-3.3 (2)			35.2 %	0.23 [0.04, 0.42]
Klaber 2000b	210	3.1 (2)	210	3.7 (2.2)	•	•	35.2 %	-0.28 [-0.48, -0.09]
Van de Kerkhof 2002a	41	-0.48 (0.24)	46	-0.17 (0.38)	-#-		29.5 %	-0.95 [-1.40, -0.51]
Subtotal (95% CI)	458		467		-	•	100.0 %	-0.30 [-0.84, 0.24]
Heterogeneity: $Tau^2 = 0.21$;	Chi ² = 29.12, df =	2 (P<0.00001)); I ² =93%					
Test for overall effect: $Z = I$.09 (P = 0.28)							
Test for subgroup difference	es: Chi ² = 143.01, d	f = 5 (P = 0.00)), I ² =97%					
					4 -2	0 2	4	
			Favours vita	amin D alone or in	combination	Favours ot	her active treatmen	t

Analysis 19.4. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 4 PAGI.

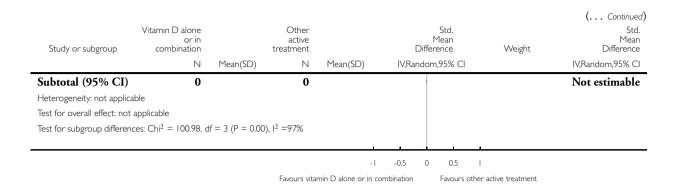
Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: 4 PAGI

Study or subgroup	Vitamin D alone or in combination		Other active treatment		Std. Mean Difference	Weight	Std Mear Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% C
I Vitamin D vs. corticoster	oid (potent): calcipo	otriol vs. BMD					
Jemec 2008 (H)	266	-3.52 (1.86)	550	-4.59 (1.39)		49.7 %	0.69 [0.54, 0.84
Van de Kerkhof 2009	282	-3.92 (1.54)	556	-4.54 (1.39)		50.3 %	0.43 [0.28, 0.57]
Subtotal (95% CI)	548		1106		-	100.0 %	0.56 [0.31, 0.81]
Heterogeneity: Tau ² = 0.03	; Chi ² = 5.80, df =	(P = 0.02); ²	=83%				
Test for overall effect: $Z = $	4.35 (P = 0.000014)					
2 Vitamin D vs. corticoster	oid (potent): calcipo	otriol vs. BMV					
Klaber 1994	236	-2.44 (1.07)	232	-2.85 (0.94)		100.0 %	0.41 [0.22, 0.59
Subtotal (95% CI)	236		232		•	100.0 %	0.41 [0.22, 0.59]
Heterogeneity: not applicat	ble						
Test for overall effect: $Z = -$	4.35 (P = 0.000014	+)					
3 Vitamin D vs. corticoster	oid (very potent): c	alcipotriol vs. cl	obetasol prop	ionate			
Subtotal (95% CI)	0		0				Not estimable
Heterogeneity: not applicat	ble						
Test for overall effect: not a	• •						
4 Vitamin D + corticosterc							
Buckley 2008	107	-4.99 (1.09)	109	-4.66 (1.28)		8.9 %	-0.28 [-0.54, -0.01
Jemec 2008 (H)	529	-4.75 (1.31)	550	-4.59 (1.39)		44.8 %	-0.12 [-0.24, 0.00
Van de Kerkhof 2009	563	-4.79 (1.23)	556	-4.54 (1.39)	-	46.3 %	-0.19 [-0.31, -0.07
Subtotal (95% CI)	1199		1215		•	100.0 %	-0.17 [-0.25, -0.09]
Heterogeneity: Tau ² = 0.0;	Chi ² = 1.43, df = 2	$2 (P = 0.49); ^2 =$	=0.0%				
Test for overall effect: $Z = $	4.06 (P = 0.000048	3)					
5 Vitamin D vs. vitamin D -	+ corticosteroid: ca	lcipotriol vs. cal	cipotriol + B№	1D			
Jemec 2008 (H)	266	-3.52 (1.86)	529	-4.75 (1.31)	-	35.6 %	0.81 [0.66, 0.96
Kragballe 2009	105	2.01 (1.14)	207	0.89 (0.9)	-	• 28.4 %	1.13 [0.88, 1.38
Van de Kerkhof 2009	282	-3.92 (1.54)	563	-4.79 (1.23)	-	36.0 %	0.65 [0.50, 0.79
Subtotal (95% CI)	653		1299		-	100.0 %	0.84 [0.61, 1.08]
Heterogeneity: $Tau^2 = 0.04$; $Chi^2 = 10.80$, df =	= 2 (P = 0.005)	; 2 =8 %				
Test for overall effect: $Z =$	6.96 (P < 0.00001)						
6 Vitamin D vs. other treat	ments: calcipotriol \	vs. coal tar polyt	herapy:				

(Continued ...)



Analysis 19.5. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 5 Combined end point (IAGI/TSS/PASI/PAGI).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: 5 Combined end point (IAGI/TSS/PASI/PAGI)

Study or subgroup	Vitamin D alone or in combination		Other active treatment		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% Cl		IV,Random,95% CI
l Vitamin D vs. corticoster	roid (potent): calcipo	otriol vs. BMD					
Jemec 2008 (H)	272	1.85 (1.15)	556	1.23 (1.07)	-	49.5 %	0.56 [0.42, 0.71]
Van de Kerkhof 2009	286	1.73 (1.1)	562	1.3 (1.07)	-	50.5 %	0.40 [0.25, 0.54]
Subtotal (95% CI)	558		1118		•	100.0 %	0.48 [0.32, 0.64]
Heterogeneity: $Tau^2 = 0.0$	I; $Chi^2 = 2.53$, $df =$	$ (P = 0.11); ^2$	=60%				
Test for overall effect: $Z =$	5.75 (P < 0.00001)						
2 Vitamin D vs. corticoster	roid (potent): calcipo	triol vs. BMV					
Duweb 2000	24	-2.63 (0.88)	18	-2.83 (0.92)		8.2 %	0.22 [-0.39, 0.83]
Klaber 1994	236	-2.51 (1.14)	232	-2.93 (1.02)	+	91.8 %	0.39 [0.20, 0.57]
Subtotal (95% CI)	260		250		•	100.0 %	0.37 [0.20, 0.55]
Heterogeneity: $Tau^2 = 0.0$;	; $Chi^2 = 0.27$, $df = 1$	$(P = 0.6 I); I^2 =$	0.0%				
Test for overall effect: Z =	4.18 (P = 0.000029))					
3 Vitamin D vs. corticoster	roid (very potent): c	alcipotriol vs. clo	betasol prop	ionate			
Reygagne 2005	75	2.36 (1.64)	76	1.76 (1.57)		100.0 %	0.37 [0.05, 0.69]
				-4	-2 0 2	4	
			Favours vita	amin D alone or in c	ombination Favours oth	ner active treatment	
							(Continued)

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Topical treatments for chronic plaque psoriasis (Review)

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V	itamin D alone or in		Other		Std. Mean		(Continue Sto Mear
Study or subgroup	combination		treatment		Difference	Weight	Difference
	Ν	Mean(SD)	Ν	Mean(SD)	IV,Random,95% CI		IV,Random,95% C
Subtotal (95% CI)	75		76		•	100.0 %	0.37 [0.05, 0.69]
Heterogeneity: not applicable							
Test for overall effect: $Z = 2.27$	· /						
4 Vitamin D + corticosteroid		•				0.0.0/	
Buckley 2008	108	0.79 (0.91)	110	1.05 (1.04)		8.9 %	-0.26 [-0.53, 0.00
Jemec 2008 (H)	541	1.09 (1.02)	556	1.23 (1.07)	•	45.0 %	-0.13 [-0.25, -0.02
Van de Kerkhof 2009	567	1.08 (1)	562	1.3 (1.07)	•	46.1 %	-0.21 [-0.33, -0.10
Subtotal (95% CI)	1216		1228		•	100.0 %	-0.18 [-0.26, -0.10
Heterogeneity: Tau ² = 0.0; Chi	² = 1.27, df = 2	$(P = 0.53); I^2 = 0$).0%				
Test for overall effect: $Z = 4.48$	8 (P < 0.00001)						
5 Vitamin D vs. vitamin D + co	orticosteroid: cal	cipotriol vs. calcip	ootriol + BM	D			
Jemec 2008 (H)	272	1.85 (1.15)	541	1.09 (1.02)	-	26.4 %	0.71 [0.56, 0.86
Kragballe 2009	105	2.2 (1.26)	207	1.17 (1.05)	-	21.2 %	0.91 [0.67, 1.16
Luger 2008	265	1.68 (1.12)	338	1.3 (1.01)	-	25.8 %	0.36 [0.20, 0.52
Van de Kerkhof 2009	286	1.73 (1.1)	567	1.08 (1)	-	26.6 %	0.63 [0.48, 0.77
Subtotal (95% CI)	928		1653		•	100.0 %	0.64 [0.44, 0.84
Heterogeneity: $Tau^2 = 0.03$; C	hi² = 16.99, df =	3 (P = 0.00071)	; I ² =82%				
Test for overall effect: Z = 6.23	B (P < 0.00001)						
6 Vitamin D vs. other treatmer	nts: calcipotriol v	s. coal tar polyth	erapy				
Barrett 2005	161	-3.48 (.)	170	-3.49 (1.17)	•	35.5 %	0.01 [-0.21, 0.22
Klaber 2000b	208	-2.77 (1.28)	209	-2.07 (1.56)	-	36.0 %	-0.49 [-0.68, -0.29
Van de Kerkhof 2002a	41	-0.48 (0.24)	46	-0.17 (0.38)	+	28.5 %	-0.95 [-1.40, -0.51
Subtotal (95% CI)	410		425		•	100.0 %	-0.45 [-0.92, 0.02
Heterogeneity: Tau ² = 0.15; C	hi ² = 19.63, df =	2 (P = 0.00005)	; l ² =90%				
Test for overall effect: $Z = 1.86$	5 (P = 0.063)						
Test for subgroup differences: ($Chi^2 = 112.96, c$	f = 5 (P = 0.00),	l ² =96%				

Analysis 19.6. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 6 Total withdrawals.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: 6 Total withdrawals

Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M- H,Random,95%	Weight	Risk Difference M- H,Random,9
	n/N	n/N	Cl		CI
I Vitamin D vs. corticosteroio	d (potent): calcipotriol vs. B	MD			
Jemec 2008 (H)	57/272	47/556	-	49.3 %	0.13 [0.07, 0.18]
Van de Kerkhof 2009	38/286	66/562	•	50.7 %	0.02 [-0.03, 0.06]
Subtotal (95% CI)	558	1118	•	100.0 %	0.07 [-0.04, 0.18]
Total events: 95 (Vitamin D al	lone or in combination), I I	3 (Other active treatme	ent)		
Heterogeneity: $Tau^2 = 0.01$; ($Chi^2 = 9.09, df = 1 (P = 0.0)$	003); I ² =89%			
Test for overall effect: $Z = 1.2$	26 (P = 0.21)				
2 Vitamin D vs. corticosteroio	d (potent): calcipotriol vs. B	MV			
Duweb 2000	0/24	0/18	+	18.3 %	0.0 [-0.09, 0.09]
Klaber 1994	20/240	9/234	-	81.7 %	0.04 [0.00, 0.09]
Subtotal (95% CI)	264	252	•	100.0 %	0.04 [0.00, 0.08]
Test for overall effect: $Z = 1.8$ 3 Vitamin D vs. corticosteroid	,		te		
Köse 1997	0/21	0/22	#	50.7 %	0.0 [-0.09, 0.09]
Köse 1997 Reygagne 2005	0/21	0/22 3/76	-	50.7 % 49.3 %	
Reygagne 2005			•	49.3 %	0.11 [0.02, 0.20]
Reygagne 2005 Subtotal (95% CI) Total events: 11 (Vitamin D al Heterogeneity: Tau ² = 0.01; C Test for overall effect: Z = 0.8	11/75 96 Ione or in combination), 3 Chi ² = 3.86, df = 1 (P = 0.0 84 (P = 0.40)	3/76 98 Other active treatment) (5); I ² =74%			0.11 [0.02, 0.20]
Reygagne 2005 Subtotal (95% CI) Total events: 11 (Vitamin D al Heterogeneity: Tau ² = 0.01; C Test for overall effect: Z = 0.6 4 Vitamin D + corticosteroid	11/75 96 Ione or in combination), 3 Chi ² = 3.86, df = 1 (P = 0.0 84 (P = 0.40)	3/76 98 Other active treatment) (5); I ² =74%		49.3 %	0.11 [0.02, 0.20] 0.05 [-0.07, 0.18]
Reygagne 2005 Subtotal (95% CI) Total events: 11 (Vitamin D al Heterogeneity: Tau ² = 0.01; C Test for overall effect: Z = 0.8	11/75 96 lone or in combination), 3 (Chi ² = 3.86, df = 1 (P = 0.0 84 (P = 0.40) vs. corticosteroid: calcipote	3/76 98 Other active treatment) (5); I ² =74% riol + BMD vs. BMD		49.3 % 100.0 %	0.0 [-0.09, 0.09] 0.11 [0.02, 0.20] 0.05 [-0.07, 0.18] 0.05 [-0.03, 0.13] 0.03 [-0.01, 0.06]
Reygagne 2005 Subtotal (95% CI) Total events: 11 (Vitamin D al Heterogeneity: Tau ² = 0.01; G Test for overall effect: Z = 0.8 4 Vitamin D + corticosteroid Buckley 2008	11/75 96 Ione or in combination), 3 Chi ² = 3.86, df = 1 (P = 0.0 84 (P = 0.40) vs. corticosteroid: calcipote 14/108	3/76 98 Other active treatment, (5); I ² =74% riol + BMD vs. BMD 9/110		49.3 % 100.0 % 21.2 %	0.11 [0.02, 0.20] 0.05 [-0.07, 0.18] 0.05 [-0.03, 0.13]
Reygagne 2005 Subtotal (95% CI) Total events: 11 (Vitamin D al Heterogeneity: Tau ² = 0.01; C Test for overall effect: Z = 0.8 4 Vitamin D + corticosteroid Buckley 2008 Jemec 2008 (H)	11/75 96 Ione or in combination), 3 (Chi ² = 3.86, df = 1 (P = 0.0 34 (P = 0.40) vs. corticosteroid: calcipote 14/108 61/541	3/76 98 Other active treatment) (5); I ² =74% riol + BMD vs. BMD 9/I I0 47/556		49.3 % 100.0 % 21.2 % 39.4 %	0.11 [0.02, 0.20] 0.05 [-0.07, 0.18] 0.05 [-0.03, 0.13] 0.03 [-0.01, 0.06]

Topical treatments for chronic plaque psoriasis (Review)

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Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M- H,Random,95%	Weight	Risk Difference M- H,Random,95%
	n/N	n/N	Cl		Cl
Jemec 2008 (H)	57/272	61/541	-	26.3 %	0.10 [0.04, 0.15]
Kragballe 2009	23/105	17/207	-	20.3 %	0.14 [0.05, 0.22]
Luger 2008	175/440	92/429	-	25.4 %	0.18 [0.12, 0.24]
Van de Kerkhof 2009	38/286	48/567	-	28.0 %	0.05 [0.00, 0.09]
Subtotal (95% CI)	1103	1744	•	100.0 %	0.11 [0.05, 0.18]
Heterogeneity: Tau ² = 0.00; CH Test for overall effect: Z = 3.45 6 Vitamin D vs. other treatmen Klaber 2000b	(P = 0.00056)	,	-	100.0 %	0.01 [-0.07, 0.09]
Subtotal (95% CI)	238	237	+	100.0 %	0.01 [-0.07, 0.09]
Total events: 72 (Vitamin D alo	ne or in combination), 69	(Other active treatmer	nt)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.27$	r (P = 0.79)				
Test for subgroup differences: C	$Chi^2 = 7.36, df = 5 (P = 0)$.20), I ² =32%			
			-1 -0.5 0 0.5 1		
	Fa	avours vitamin D alone or ir	n combination Favours other	active treatment	

Analysis 19.7. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 7 Withdrawals due to adverse events.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: 7 Withdrawals due to adverse events

Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Vitamin D vs. corticosteroi	d (potent): calcipotriol vs. B	MD			
Jemec 2008 (H)	20/272	6/556	•	47.1 %	0.06 [0.03, 0.09]
Van de Kerkhof 2009	8/286	7/562	-	52.9 %	0.02 [-0.01, 0.04]
Subtotal (95% CI)	558	1118	•	100.0 %	0.04 [-0.01, 0.09]
Total events: 28 (Vitamin D a	lone or in combination), 13	(Other active treatmen	t)		
Heterogeneity: Tau ² = 0.00; 0	$Chi^2 = 6.74, df = 1 (P = 0.0)$	I); I ² =85%			
Test for overall effect: $Z = 1.4$	48 (P = 0.14)				
2 Vitamin D vs. corticosteroi	d (potent): calcipotriol vs. B	MV			
Duweb 2000	0/24	0/18	+	9.3 %	0.0 [-0.09, 0.09]
Klaber 1994	1/240	2/234	•	90.7 %	0.04 [0.01, 0.07]
Subtotal (95% CI)	264	252	•	100.0 %	0.03 [0.01, 0.06]
Test for overall effect: Z = 2.4 3 Vitamin D vs. corticosteroid Köse 1997	· /	vs. clobetasol propionat 0/22	e 🖷	46.5 %	0.0 [-0.09, 0.09]
Reygagne 2005	7/75	0/76	-	53.5 %	0.09 [0.02, 0.16]
They gagne 2000	,,,,,,	98		100.0 %	2 3
Subtatal (95% CI)	96				0 05 1 -0 05 0 15 1
Subtotal (95% CI) Total events: 7 (Vitamin D ald Heterogeneity: Tau ² = 0.00; 0 Test for overall effect: Z = 1.0 4 Vitamin D + corticosteroid	$Chi^2 = 3.07, df = 1 (P = 0.0)$	Other active treatment) 8); $l^2 = 67\%$		100.0 /0	0.05 [-0.05, 0.15]
Total events: 7 (Vitamin D ald Heterogeneity: Tau ² = 0.00; $($	one or in combination), 0 (C Chi ² = 3.07, df = 1 (P = 0.0 D1 (P = 0.31)	Other active treatment) 8); $l^2 = 67\%$		7.4 %	
Total events: 7 (Vitamin D alo Heterogeneity: Tau ² = 0.00; 0 Test for overall effect: Z = 1.0 4 Vitamin D + corticosteroid	one or in combination), 0 (C Chi ² = 3.07, df = 1 (P = 0.0 01 (P = 0.31) 1 vs. corticosteroid: calcipot	Dther active treatment) (8); I ² =67% riol + BMD vs. BMD			0.05 [-0.05, 0.15] -0.01 [-0.04, 0.02] 0.00 [-0.01, 0.02]
Total events: 7 (Vitamin D alc Heterogeneity: Tau ² = 0.00; 0 Test for overall effect: Z = 1.0 4 Vitamin D + corticosteroid Buckley 2008	cone or in combination), 0 (C Chi ² = 3.07, df = 1 (P = 0.0 D1 (P = 0.31) I vs. corticosteroid: calcipotr 1/108	Dther active treatment) 8); I ² =67% iol + BMD vs. BMD 2/110		7.4 %	-0.01 [-0.04, 0.02]
Total events: 7 (Vitamin D alc Heterogeneity: Tau ² = 0.00; (Test for overall effect: Z = 1.0 4 Vitamin D + corticosteroid Buckley 2008 Jemec 2008 (H)	cone or in combination), 0 (C Chi ² = 3.07, df = 1 (P = 0.0 D1 (P = 0.31) I vs. corticosteroid: calcipotr 1/108 8/541	Dther active treatment) 8); I ² =67% iol + BMD vs. BMD 2/110 6/556		7.4 % 39.5 %	-0.01 [-0.04, 0.02] 0.00 [-0.01, 0.02]

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Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M- H,Random,95%	Weight	Risk Difference H,Random,95%
	n/N	n/N	Cl		Cl
Jemec 2008 (H)	20/272	8/541	•	25.7 %	0.06 [0.03, 0.09]
Kragballe 2009	9/105	2/207	-	18.2 %	0.08 [0.02, 0.13]
Luger 2008	44/440	9/429	-	26.3 %	0.08 [0.05, 0.11]
Van de Kerkhof 2009	8/286	4/567	-	29.8 %	0.02 [0.00, 0.04]
Subtotal (95% CI)	1103	1744	*	100.0 %	0.06 [0.02, 0.09]
Total events: 81 (Vitamin D ald Heterogeneity: Tau ² = 0.00; C Test for overall effect: Z = 3.12 6 Vitamin D vs. other treatmen Klaber 2000b	$hi^2 = 14.23, df = 3 (P = 0)$ 3 (P = 0.0017)	003); I ² =79%	••)	100.0 %	0.08 [0.02, 0.14]
Subtotal (95% CI)	230	215	•	100.0 %	0.08 [0.02, 0.14]
Total events: 35 (Vitamin D alo	one or in combination), 16	(Other active treatment	nt)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 2.62$	2 (P = 0.0088)				
Test for subgroup differences:	Chi ² = 23.39, df = 5 (P =	0.00), l ² =79%			
			-1 -0.5 0 0.5 1		
	Fa	avours vitamin D alone or i	n combination Favours other	active treatment	

Analysis 19.8. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 8 Withdrawals due to treatment failure.

Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: 8 Withdrawals due to treatment failure

Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Vitamin D vs. corticosteroio	d (potent): calcipotriol vs. E	MD			
Jemec 2008 (H)	19/272	9/556	•	46.4 %	0.05 [0.02, 0.09]
Van de Kerkhof 2009	8/286	9/562	-	53.6 %	0.01 [-0.01, 0.03]
Subtotal (95% CI)	558	1118	•	100.0 %	0.03 [-0.01, 0.07]
Total events: 27 (Vitamin D al	one or in combination), 18	(Other active treatmer	nt)		
Heterogeneity: $Tau^2 = 0.00$; ($Chi^2 = 5.09, df = 1 (P = 0.0)$	02); I ² =80%			
Test for overall effect: Z = 1.4	+I (P = 0.16)				
2 Vitamin D vs. corticosteroio	d (potent): calcipotriol vs. E	BMV			
Duweb 2000	0/24	0/18		4.7 %	0.0 [-0.09, 0.09]
Klaber 1994	4/240	2/234	•	95.3 %	0.01 [-0.01, 0.03]
Subtotal (95% CI)	264	252	•	100.0 %	0.01 [-0.01, 0.03]
Test for overall effect: Z = 0.7 3 Vitamin D vs. corticosteroic Köse 1997	· /	vs. clobetasol propiona 0/22	te	14.9 %	0.0 [-0.09, 0.09]
			-	14.9 %	0.0 [-0.09, 0.09]
Reygagne 2005	1/75	0/76	-	85.1 %	0.01 [-0.02, 0.05]
Subtotal (95% CI)	96	98	+	100.0 %	0.01 [-0.02, 0.04]
Total events: (Vitamin D alo Heterogeneity: Tau ² = 0.0; Cł Test for overall effect: Z = 0.6 4 Vitamin D + corticosteroid	$hi^2 = 0.08$, df = 1 (P = 0.78) 57 (P = 0.50)	3); I ² =0.0%			
Buckley 2008	0/108	2/110	-	7.9 %	-0.02 [-0.05, 0.01]
Jemec 2008 (H)	2/541	9/556	•	53.7 %	-0.01 [-0.02, 0.00]
Van de Kerkhof 2009	7/567	9/562	-	38.4 %	0.00 [-0.02, 0.01]
Subtotal (95% CI)	1216	1228	•	100.0 %	-0.01 [-0.02, 0.00]
Total events: 9 (Vitamin D alo	ne or in combination), 20	(Other active treatment)		
Heterogeneity: $Tau^2 = 0.0$; Ch	$hi^2 = 1.27, df = 2 (P = 0.52)$	3); l ² =0.0%			
Test for overall effect: $Z = 2.1$	9 (P = 0.029)				
5 Vitamin D vs. vitamin D + c	orticosteroid: calcipotriol	vs. calcipotriol + BMD			
5 Vitamin D vs. vitamin D + c	corticosteroid: calcipotriol		1.5 -0.25 0 0.25 0.5 combination Favours other	active treatment	

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Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference H,Random,95%	Weight	Risk Difference M- H,Random,95%
	n/N	n/N	Cl		Cl
Jemec 2008 (H)	19/272	2/541	-	32.8 %	0.07 [0.04, 0.10]
Luger 2008	51/440	14/429	-	31.8 %	0.08 [0.05, 0.12]
Van de Kerkhof 2009	8/286	7/567	-	35.4 %	0.02 [-0.01, 0.04]
Subtotal (95% CI)	998	1537	•	100.0 %	0.05 [0.01, 0.10]
Total events: 78 (Vitamin D alo	one or in combination), 23	(Other active treatmen	it)		
Heterogeneity: $Tau^2 = 0.00$; C	$Chi^2 = 16.89, df = 2 (P = 0)$	0.00021); I ² =88%			
Test for overall effect: $Z = 2.2$	I (P = 0.027)				
6 Vitamin D vs. other treatme	nts: calcipotriol vs. coal tar	polytherapy			
Subtotal (95% CI)	0	0			Not estimable
Total events: 0 (Vitamin D alor	ne or in combination), 0 (0	Other active treatment)			
Heterogeneity: not applicable					
Test for overall effect: not app	licable				
Test for subgroup differences:	Chi ² = 11.94, df = 4 (P =	0.02), I ² =66%			
				1	
		-0	.5 -0.25 0 0.25	0.5	
	Fa	wours vitamin D alone or in	combination Favours oth	ner active treatment	

Analysis 19.9. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 9 Adverse events (local).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: 9 Adverse events (local)

Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,S Cl
I Vitamin D vs. corticosteroid	(potent): calcipotriol vs. E	MD			
Jemec 2008 (H)	35/266	29/548	-	48.6 %	0.08 [0.03, 0.12]
Van de Kerkhof 2009	36/282	32/556	-	51.4 %	0.07 [0.03, 0.11]
Subtotal (95% CI)	548	1104	•	100.0 %	0.07 [0.04, 0.11]
Total events: 71 (Vitamin D alo	one or in combination), 6 l	(Other active treatmen	t)		
Heterogeneity: $Tau^2 = 0.0$; Ch	$i^2 = 0.07$, df = 1 (P = 0.79)	9); I ² =0.0%			
Test for overall effect: $Z = 4.67$	7 (P < 0.00001)				
2 Vitamin D vs. corticosteroid	(potent): calcipotriol vs. E	MV			
Duweb 2000	2/24	0/18	-	43.3 %	0.08 [-0.05, 0.22]
Klaber 1994	84/240	26/234	-	56.7 %	0.24 [0.17, 0.31]
Subtotal (95% CI)	264	252	•	100.0 %	0.17 [0.01, 0.33]
Total events: 86 (Vitamin D ald Heterogeneity: Tau ² = 0.01; C Test for overall effect: Z = 2.1	$hi^2 = 4.25, df = 1 (P = 0.0)$		t)		
3 Vitamin D vs. corticosteroid	· /	vs. clobetasol propionat	e		
Köse 1997	4/21	2/22		18.4 %	0.10 [-0.11, 0.31
Reygagne 2005	17/75	1/76	-	81.6 %	0.21 [0.12, 0.31]
Subtotal (95% CI)	96	98	•	100.0 %	0.19 [0.10, 0.28]
Total events: 21 (Vitamin D alo Heterogeneity: Tau ² = 0.0; Ch Test for overall effect: Z = 4.20 4 Vitamin D + corticosteroid	$i^2 = 0.96$, df = 1 (P = 0.32) 6 (P = 0.000021)	8); ² =0.0%)		
Buckley 2008	1/108	2/110	-	27.5 %	-0.01 [-0.04, 0.02
Jemec 2008 (H)	25/530	29/548	-	38.6 %	-0.01 [-0.03, 0.02]
Van de Kerkhof 2009	35/563	32/556	+	33.8 %	0.00 [-0.02, 0.03]
Subtotal (95% CI)	1201	1214		100.0 %	0.00 [-0.02, 0.01]
otal events: 61 (Vitamin D alo	one or in combination), 63 $i^2 = 0.52$, df = 2 (P = 0.72)		t)		

(Continued \dots)

(... Continued)

Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
5 Vitamin D vs. vitamin D + c	orticosteroid: calcipotriol	vs. calcipotriol + BMD			
Jemec 2008 (H)	35/266	25/530	•	30.4 %	0.08 [0.04, 0.13]
Kragballe 2009	20/104	7/206	+	12.4 %	0.16 [0.08, 0.24]
Luger 2008	93/431	50/419	-	26.1 %	0.10 [0.05, 0.15]
Van de Kerkhof 2009	36/282	35/563	-	31.1 %	0.07 [0.02, 0.11]
Subtotal (95% CI)	1083	1718	•	100.0 %	0.09 [0.06, 0.12]
Total events: 184 (Vitamin D a	alone or in combination), I	17 (Other active treatm	ient)		
Heterogeneity: $Tau^2 = 0.00$; C	$Chi^2 = 4.17, df = 3 (P = 0.2)$	24); I ² =28%			
Test for overall effect: $Z = 5.8$	9 (P < 0.00001)				
6 Vitamin D vs. other treatme	nts: calcipotriol vs. coal tar	polytherapy			
Klaber 2000b	137/230	77/215		100.0 %	0.24 [0.15, 0.33]
Subtotal (95% CI)	230	215	•	100.0 %	0.24 [0.15, 0.33]
Total events: 137 (Vitamin D a	alone or in combination), 7	7 (Other active treatme	nt)		
Heterogeneity: not applicable					
Test for overall effect: $Z = 5.1$	6 (P < 0.00001)				
Test for subgroup differences:	Chi ² = 74.84, df = 5 (P =	0.00), I ² =93%			
		-	-I -0.5 0 0.5 I		
	F	avours vitamin D alone or in	combination Favours other	active treatment	

Analysis 19.10. Comparison 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments, Outcome 10 Adverse events (systemic).

Review: Topical treatments for chronic plaque psoriasis

Comparison: 19 Scalp psoriasis: vitamin D alone or in combination versus other treatments

Outcome: 10 Adverse events (systemic)

Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M-	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,9 Cl
I Vitamin D vs. corticosteroi	d (potent): calcipotriol vs. B	MD			
Jemec 2008 (H)	0/272	0/556	-	48.6 %	0.0 [-0.01, 0.01]
Van de Kerkhof 2009	0/282	0/556	-	51.4 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	554	1112		100.0 %	0.0 [0.00, 0.00]
Total events: 0 (Vitamin D ald Heterogeneity: Tau ² = 0.0; C Test for overall effect: Z = 0.0	$hi^2 = 0.0, df = 1 (P = 1.00);$ 0 (P = 1.0)	; ² =0.0%			
2 Vitamin D vs. corticosteroi Klaber 1994	d (potent): calcipotriol vs. Bl 0/240	0/234		100.0 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	240	234	Ţ	100.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Vitamin D alc Heterogeneity: not applicable Test for overall effect: Z = 0.0 3 Vitamin D vs. corticosteroio	e D (P = 1.0)		2		
Reygagne 2005	0/75	0/76	-	100.0 %	0.0 [-0.03, 0.03]
Subtotal (95% CI)	75	76	+	100.0 %	0.0 [-0.03, 0.03]
Fotal events: 0 (Vitamin D ald Heterogeneity: not applicable Fest for overall effect: Z = 0.0 4 Vitamin D + corticosteroid Jemec 2008 (H)	e D (P = 1.0)		_	49.0 %	0.0 [0.00, 0.00]
Van de Kerkhof 2009	0/563	0/556		51.0 %	0.0 [0.00, 0.00]
Subtotal (95% CI)	1104	1112		100.0 %	0.0 [0.00, 0.00]
Total events: 0 (Vitamin D alc	, (,			
Heterogeneity: $Tau^2 = 0.0$; C Test for overall effect: $Z = 0.0$	D (P = 1.0)	: calcinotrial + BMD			
8 ,	D (P = 1.0)	s. calcipotriol + BMD 0/541	-	44.9 %	0.0 [-0.01, 0.01]
Test for overall effect: Z = 0.0 5 Vitamin D vs. vitamin D + 0	P(P = 1.0)		-	44.9 % 6.8 %	0.0 [-0.01, 0.01]

(Continued ...)

(... Continued)

Study or subgroup	Vitamin D alone or in combination	Other active treatment	Risk Difference M	Weight	Risk Difference M-
	n/N	n/N	H,Random,95% Cl		H,Random,95% Cl
Subtotal (95% CI)	659	1311		100.0 %	0.0 [0.00, 0.00]
Total events: 0 (Vitamin D alor	ne or in combination), 0 (C	ther active treatment)			
Heterogeneity: $Tau^2 = 0.0$; Ch	i ² = 0.0, df = 2 (P = 1.00);	$ ^2 = 0.0\%$			
Test for overall effect: $Z = 0.0$	(P = 1.0)				
6 Vitamin D vs. other treatme	nts: calcipotriol vs. coal tar	polytherapy			
Klaber 2000b	0/230	0/215	-	100.0 %	0.0 [-0.01, 0.01]
Subtotal (95% CI)	230	215	+	100.0 %	0.0 [-0.01, 0.01]
Total events: 0 (Vitamin D alor	ne or in combination), 0 (C	ther active treatment)			
Heterogeneity: not applicable					
Test for overall effect: $Z = 0.0$	(P = 1.0)				
Test for subgroup differences:	Chi ² = 0.0, df = 5 (P = 1.0	0), l ² =0.0%			
		-(0.2 -0.1 0 0.1	0.2	
	F	n de la compansión de la c			

Favours vitamin D alone or in combination Favours other active treatment

ADDITIONAL TABLES

Table 1. List of acronyms

Acronym	Full name
BC	baseline comparability demonstrated (clinical/demographic)
BD	twice daily
BMD	betamethasone dipropionate
BMV	betamethasone valerate
BSA	Body Surface Area
Btw-patient	Between-patient
CI	confidence interval
dys	days
EQ-5D	EuroQOL
FU	follow up (includes treatment period)

Table 1. List of acronyms (Continued)

I ²	heterogeneity statistic
IAGI	Investigator Assessment of Global Improvement (change score)
IGA	Investigator Global Assessment (static score)
IQR	interquartile range
ISGA	Investigator's Static Global Assessment Score
LAE	local adverse effects
LCD	liquor carbonis distillate
LF	loss to follow up (per cent of participants randomised, not contributing to primary outcome measure)
MEMS	Medication Event Monitoring System
mPASI	modified Psoriasis Area Severity Index
NA	not available/not applicable
NR	not reported
OD	once daily
ОМ	once in the morning
ON	once at night
ODS	overall disease severity
PAGI	Patient Assessment of Global Improvement (change score)
PASI	Psoriasis Area Severity Index
PDI	Psoriasis Disability Index
PGA	Patient Global Assessment (static score)
PMAQ-3w	Medication Adherence Questionnaire, version 3W
pt	point
QOL	quality of life
RD	risk difference

SD	standard deviation				
SMD	standardised mean difference				
ТСР	two-compound product				
TD	three times daily				
TLPSS	Total Local Psoriasis Severity Score				
TSS	Total Severity Score/total sum score				
UV	ultra violet				
VDRE	Vitamin D-Responsive Element				
wks	weeks				
yrs	years				

Table 2. Overview of outcome measures on effectiveness

Outcome	Acronym	Construct	Scale, minimum	Scale, maximum	Notes
* Investigator's As- sessment of Over- all Global Improve- ment	IAGI	Improvement from baseline variably defined. Common tax- onomy ranges from worse to cleared	4-pt	7-pt	Calculated means and standard deviations by assigning zero to 'worse' (or equivalent). Higher scores indicate greater improvement
Inves- tigator's Global As- sessment of Disease Severity	IGA	Static equivalent of the IAGI	4-pt	7-pt	Calculated means and standard deviations by assigning zero to 'clear' (or equivalent). Higher scores indicate more severe disease
Total Severity Score	TSS	Redness (ery- thema), thickness (in- filtration) and scaling (sometimes also itch- ing (pruritis)) of target plaque(s). Scored sep- arately then summed	0 to 3	0 to 24	Also known as the Lo- cal Psoriasis Severity Index or the Total Sum Score. Higher scores indicate more severe disease

Table 2. Overview of outcome measures on effectiveness (Continued)

Psoriasis Area and Severity Index	PASI	Redness, thickness, and scaliness of the le- sions (each graded on a 0 to 4 scale), weighted by the area of involve- ment (0 to 6) and summed	0 to 68 (without head)	0 to 72 (including head)	Higher scores indicate more severe disease
* Patient's Assessment of Over- all Global Improve- ment	PAGI	Assessed as IAGI	4-pt	7-pt	Less often reported than IAGI. Majority of included trials use 5-pt scale
Patient's Global As- sessment of Disease Severity	PGA	Assessed as IGA	4-pt	5-pt	Rarely reported (5/ 177 studies)

* IAGI/PAGI data are entered as a negative values; thus, a reduction denotes a positive improvement for the active treatment consistent with TSS and PASI measures

Type of study/score	Placebo IAGI (change) /IGA (end point)	Placebo TSS	Placebo PASI	Placebo PAGI (change) /PGA (end point)	H2H IAGI (change) /IGA (end point)	H2H TSS	H2H PASI	H2H PAGI (change)/ PGA (end point)
Between- patient (end point)	0.93	1.33	3.76	1.13	1.01	1.65	3.61	1.12
Within- patient (end point)	1.08	1.49	7.17	NA	NA	1.50	2.58	NA
Between- patient (change)	1.17	1.52	5.75	1.31	1.10	1.73	7.85	1.20
Within- patient (change)	1.02	1.58	NA	1.53	0.96	1.94	NA	0.83
Within- patient (% change)	NA	0.18	NA	NA	NA	NA	NA	NA

Table 3. Summary of imputed standard deviation values

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Table 3.	Summary of imputed standard deviation values	(Continued)
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Between- patient (% change)	NA	NA	0.37	NA	NA	0.13	0.33	NA
Scalp between- patient (end point)	1.08	1.74	NA	1.06	1.06	1.94	NA	1.18
Scalp within- patient (end point)	1.33	NA						
Scalp between- patient (change)	1.20	NA	NA	1.28	1.30	1.75	NA	1.20
Scalp between- patient (% change)	NA	NA	NA	NA	NA	0.25	NA	NA

NA: not available; H2H: head-to-head; IGA [PGA]: Investigator [Patient] Global Assessment of Disease Severity; IAGI [PAGI]: Investigator (patient) Assessment of Global Improvement; TSS: Total Severity Score; PASI: Psoriasis Area and Severity Index

Comparison No.	Comparison Label	No. studies (NB: a study may con- tribute to more than one compari- son)	Per cent studies with between-patient design	No. participants
01	Vitamin D analogues vs. placebo	30	60%	4986
02	Corticosteroid (potent) vs. placebo	13	85%	2216
03	Corticosteroid (very po- tent) vs. placebo	10	70%	1264
04	Dithranol vs. placebo	3	0%	47

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05	Vitamin D combination products vs. placebo	5	100%	2058
06	Other treatment vs. placebo	26	46%	1450
07	Vitamin D analogues vs. corticosteroid (potent)	14	64%	3542
08	Vitamin D analogues vs. corticosteroid (very po- tent)	2	100%	82
09	Vitamin D combined with corticosteroid vs. corticos- teroid	5	100%	2113
10	Vitamin D alone or in combination vs. dithranol	8	88%	1284
11	Vitamin D alone or in combination vs. other vi- tamin D analogue	4	75%	513
12	Vitamin D alone or in combination vs. vitamin D + corticosteroid	17	94%	5856
13	Vitamin D alone or in combination vs. other treatments: com- plex regimens	9	89%	2936
14	Vitamin D alone or in combination vs. other treatment: long- term studies (> 24 wks)	1	100%	297
15	Vitamin D analogues vs. other treatment	19	68%	2364
16	Flexural/facial psoriasis: placebo-controlled trials	2	100%	122
17	Flexural/facial pso- riasis: vitamin D alone or in combination vs. other treatment	4	75%	588

Table 4. Overview of analyses: evidence of effectiveness outcomes (Continued)

18	Scalp psoriasis: placebo- controlled trials	14	93%	3011
19	Scalp psoriasis: vitamin D alone or in combination vs. other treatments	12	100%	5413

Table 4. Overview of analyses: evidence of effectiveness outcomes (Continued)

Table 5. Analysis 01: Trial characteristics and outcomes: vitamin D vs. placebo

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Calcipotriol OD/BD	Effect size [CI]	(SMD -0.93; 95% CI -1.17 to -0.68)	(SMD -1.15; 95% CI -1.41 to -0.89)	(SMD -0.65; 95% CI -0.75 to -0.55)	(SMD -0.64; 95% CI -0.97 to -0.30)	(SMD -0.96; 95% CI -1.15 to -0.77)
02 Calcipotriol plus occlusion	Effect size [CI]	NA	(SMD -0.15; 95% CI -0.44 to 0.14)	-	-	(SMD -0.15; 95% CI -0.44 to 0.14)
03 Calcitriol OD/BD	Effect size [CI]	(SMD -1.03; 95% CI -1.71 to -0.36)	(SMD -1.22; 95% CI -2.38 to -0.07)	-	(SMD -0.59; 95% CI -0.76 to -0.41)	(SMD -0.92; 95% CI -1.54 to -0.29)
04 Tacalcitol OD	Effect size [CI]	(SMD -0.84; 95% CI -1.41 to -0.26)	· · · · · ·	(SMD -0.27; 95% CI -0.56 to 0.03)	(SMD -0.24; 95% CI -0.53 to 0.05)	(SMD -0.73; 95% CI -1.09 to -0.37)
05 Maxacalcitol OD	Effect size [CI]	(SMD -1.43; 95% CI -1.91 to -0.96)	(SMD -1.61; 95% CI -2.10 to -1.12)	-	-	(SMD -1.43; 95% CI -1.91 to -0.96)
06 Paricalcitol OD	Effect size [CI]	(SMD -1.66; 95% CI -2.66 to -0.67)	(SMD -2.15; 95% CI -3.24 to -1.06)	-	-	(SMD -1.66; 95% CI -2.66 to -0.67)
07 Becocalcidiol OD	Effect size [CI]	(SMD -0.22; 95% CI -0.58 to 0.14)	(SMD -0.02; 95% CI -0.37 to 0.34)	-	-	(SMD -0.22; 95% CI -0.58 to 0.14)
08 Becocalcidiol BD	Effect size [CI]	(SMD -0.67; 95% CI -1.04 to -0.30)	(SMD -0.46; 95% CI -0.83 to -0.10)	-	-	(SMD -0.67; 95% CI -1.04 to -0.30)
All treatments	Effect size [CI]; I ² statistic		(SMD -1.04; 95% CI - 1.33 to -0.74) I ² statistic: 93.0%		(SMD -0.54; 95% CI -0.72 to -0.36):	(SMD -0.90; 95% CI -1.07 to -0.72);

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		0%		I ² statistic: 42. 3%	I ² statistic: 55. 5%	I ² statistic: 87. 5%
-	No. participants	3771	2647	2357	1467	4986
-	Between-patient design	13	9	8	5	18
-	Within-patient design	7	10	1	0	12
-	Treatment dura- tion	4 wks to 12 wks	4 wks to 12 wks	3 wks to 8 wks	8 wks to 8 wks	3 wks to 12 wks
Sensitivity analy- ses	Within-patient trials	-	-	-	-	(SMD -1.11; 95% CI -1.58 to -0.64)
-	Between-patient trials	-	-	-	-	(SMD -0.80; 95% CI -0.96 to -0.63)
-	Calcitriol, Perez 1996 removed	-	-	-		(SMD -0.60; 95% CI -0.78 to -0.41)
-	Calcipotriol BD	-	-	-	-	(SMD -1.02; 95% CI -1.23 to -0.82)
-	Calcipotriol OD	-	-	-	-	(SMD -0.76; 95% CI -1.13 to -0.40)
-	correlation coef- ficient (rho) = 0 All trials	-	-	-	-	(SMD -0.85; 95% CI -1.00 to -0.71); I ² statistic: 87. 8%
-	rho = 0 Btw-patient tri- als rho = 0.25 Within-patient trials	-	-	-	-	(SMD -0.87; 95% CI -1.01 to -0.72); I ² statistic: 88. 8%
-	rho = 0 Btw-patient tri-	-	-	-	-	(SMD -0.88; 95% CI -1.03 to

Table 5. Analysis 01: Trial characteristics and outcomes: vitamin D vs. placebo (Continued)

Table 5.	Analysis 01: Trial characteristics and outcom	es: vitamin D vs. placebo	(Continued)
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als rho = 0.50 Within-patier trials	nt				-0.73); I ² statistic = 90. 3%
- rho = 0 Btw-patient als rho = 0.75 Within-patien trials		-	-	-	(SMD -0.91; 95% CI -1.07 to -0.75); I ² statistic: 93. 2%

Table 6. Analysis 02: Trial characteristics and outcomes: potent steroids vs. placebo

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Betametha- sone dipropi- onate OD	Effect size [CI]	· · · · · · · · · · · · · · · · · · ·	(SMD -0.74; 95% CI -1.16 to -0.32)	(SMD -0.79; 95% CI -1.44 to -0.14)	-	(SMD -0. 80; 95% CI -0.96 to -0.64)
02 Betametha- sone dipropi- onate BD	Effect size [CI]		(SMD -0.77; 95% CI -1.48 to -0.06)	(SMD -1.21; 95% CI -1.44 to -0.97)	-	(SMD -1. 35; 95% CI -1.56 to -1.15)
03 Betametha- sone dipro- pionate, mainte- nance	Effect size [CI]		(SMD -0.95; 95% CI -1.62 to -0.27)	-	-	(SMD -0. 95; 95% CI -1.62 to -0.27)
04 Betametha- sone valerate	Effect size [CI]	(SMD -1.41; 95% CI -1.92 to -0.90)	(SMD -1.09; 95% CI -2.00 to -0.18)	-	-	(SMD -1. 33; 95% CI -1.78 to -0.89)
05 Budesonide	Effect size [CI]	-	-	-	-	-
06 Desonide	Effect size [CI]	· · · · · · · · · · · · · · · · · · ·	(SMD -1.16; 95% CI -1.70 to -0.61)	-	-	(SMD -0. 81; 95% CI -1.34 to -0.28)
07 Diflorasone diacetate	Effect size [CI]	-	(SMD -0.32; 95% CI -0.73 to 0.09)	-	-	(SMD -0. 32; 95% CI -0.73 to 0.09)

Table 6.	Analysis 02: Ti	rial characteristics and	outcomes: potent steroids vs.	placebo	(Continued)
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08 Fluticasone propionate	Effect size [CI]	(SMD -0.93; 95% CI -1.14 to -0.72)	-	-	-	(SMD -0. 93; 95% CI -1.14 to -0.72)
09 Hydrocorti- sone buteprate	Effect size [CI]	-	(SMD -0.46; 95% CI -0.77 to -0.15)	-	-	(SMD -0. 46; 95% CI -0.77 to -0.15)
10 Mometasone furoate	Effect size [CI]	(SMD -0.75; 95% CI -1.17 to -0.34)	(SMD -1.12; 95% CI -1.55 to -0.68)	-	-	(SMD -0. 75; 95% CI -1.17 to -0.34)
All treatments	Effect size [CI]; I ² statistic	(SMD -1.00; 95% CI - 1.18 to -0.82); I ² statistic: 57.6%		(SMD -0.97; 95% CI - 1.31 to -0.62); I ² statistic: 79.6%	-	(SMD -0.89; 95% CI -1.06 to - 0.72); I ² statistic: 65.1%
-	No. participants	1867	553	1158	0	2216
-	Between-patient design	8	6	3	0	11
-	Within-patient design	1	1	0	0	2
-	Treatment dura- tion	3 wks to 12 wks	2 wks to 12 wks	4 wks to 8 wks	-	2 wks to 12 wks
Sensitivity analy- ses	Within-patient trials	-	-	-	-	(SMD -1. 33; 95% CI -1.78 to -0.89)
-	Between-patient trials	-	-	-	-	(SMD -0. 85; 95% CI -1.03 to -0.67)
-	correlation coef- ficient (rho) = 0 All trials	-	-	-	-	(SMD -0. 89; 95% CI -1.06 to -0.72) I ² statis- tic: 77.7%
-	rho = 0 Btw-patient tri- als rho = 0.25 Within-patient trials	-	-	-	-	(SMD -0. 89; 95% CI -1.06 to -0.72) I ² statis- tic: 78.0%

 Table 6. Analysis 02: Trial characteristics and outcomes: potent steroids vs. placebo
 (Continued)

-	rho = 0 Btw-patient tri- als rho = 0.50 Within-patient trials	-	-	-	-	(SMD -0. 90; 95% CI -1.07 to -0.73) I ² statis- tic: 78.6%
-	rho = 0 Btw-patient tri- als rho = 0.75 Within-patient trials	-	-	-	-	(SMD -0. 91; 95% CI -1.08 to -0.74) I ² statis- tic: 80.2%

For acronyms, see Table 1. Both within-patient trials compared betamethasone valerate with placebo.

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Clobetasol propionate	Effect size [CI]		(SMD -1.35; 95% CI -1.80 to -0.89)	-	(SMD -1.01; 95% CI -1.55 to -0.47)	(SMD -1.65; 95% CI -2.10 to -1.20)
02 Halcinonide	Effect size [CI]	-	-	-	-	-
03 Halobetasol	Effect size [CI]	(SMD -1.81; 95% CI -2.37 to -1.24)	-	-	(SMD -1.25; 95% CI -1.46 to -1.04)	(SMD -1.36; 95% CI -1.65 to -1.07)
All treatments	Effect size [CI], N, I ²	-1.87; 95% CI -	(SMD -1.35; 95% CI - 1.80 to -0.89); I ² statistic: 75.3%	-	(SMD -1.22; 95% CI -1.42 to -1.02); I ² statistic: 0%	(SMD -1.56; 95% CI -1.87 to -1.26); I ² statistic: 81.7%
-	No. participants	515	545	0	283	1264
-	Between-patient design	4	3	0	1	7
-	Within-patient design	1	0	0	2	3
-	Treatment dura- tion	2 wks to 4 wks	2 wks to 4 wks		2 wks to 2 wks	2 wks to 4 wks

Table 7.	Analysis 03:	Trial characteristics and	outcomes: v. p	ootent steroids vs.	placebo

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Sensitivity analy- ses	Within-patient trials	-	-	-	-	(SMD -1.52; 95% CI -2.02 to -1.02)
-	Between-patient trials	-	-	-	-	(SMD -1.58; 95% CI -1.99 to -1.17)
-	correlation coef- ficient (rho) = 0 All trials	-	-	-	-	(SMD -1.52; 95% CI -1.80 to -1.24) I ² statistic: 81.6%
-	rho = 0 Btw-patient tri- als rho = 0.25 Within-patient trials	-	-	-	-	(SMD -1.52; 95% CI -1.80 to -1.25) I ² statistic: 82.2%
-	rho = 0 Btw-patient tri- als rho = 0.50 Within-patient trials	-	-	-	-	(SMD -1.53; 95% CI -1.80 to -1.26) I ² statistic: 83.3%
-	rho = 0 Btw-patient tri- als rho = 0.75 Within-patient trials	-	-	-	-	(SMD -1.55; 95% CI -1.80 to -1.29) I ² statistic: 85.9%

Table 7. Analysis 03: Trial characteristics and outcomes: v. potent steroids vs. placebo (Continued)

Table 8. Analysis 05: Trial characteristics and outcomes: vitamin D combination vs. placebo

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Combination calcipotriol/ betamethasone dipropionate, OD	Effect size [CI]	(SMD -1.21; 95% CI -1.50 to -0.91)	-	X	(SMD -0.69; 95% CI -0.98 to -0.40)	(SMD -1.21; 95% CI -1.50 to -0.91)
02 Combination calcipotriol/ betamethasone dipropionate,	Effect size [CI]	(SMD -1.90; 95% CI -2.09 to -1.71)	-	(SMD -1.41; 95% CI -1.86 to -0.97)	-	(SMD -1.90; 95% CI -2.09 to -1.71)

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BD						
All treatments	Effect size [CI], N, I ² statistic	(SMD -1.44; 95% CI - 1.76 to -1.12); I ² statistic: 89.4%	-	CI -1.53 to -0.95);	(SMD -0.69; 95% CI -0.98 to -0.40); I ² statistic: NA	(SMD -1.44; 95% CI -1.76 to -1.12); I ² statistic: 89.4%
-	No. participants	2058	0	2056	235	2058
-	Between-patient design	5	0	5	1	5
-	Within-patient design	0	0	0	0	0
- For acronyms, see	Treatment dura- tion	4 wks to 8 wks	-	4 wks to 8 wks	8 wks	4 wks to 8 wks

 Table 8. Analysis 05: Trial characteristics and outcomes: vitamin D combination vs. placebo
 (Continued)

Table 9. Analysis 06: Trial characteristics and outcomes: other treatments vs. placebo

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Aloe vera ex- tract	Effect size [CI]	-	-	(SMD -1.58; 95% CI -2.16 to -0.99)	-	(SMD -1.58; 95% CI -2.16 to -0.99)
02 Anti- IL-8 monoclonal antibody cream	Effect size [CI]	(SMD -0.59; 95% CI -1.01 to -0.16)	(SMD -0.70; 95% CI -1.13 to -0.27)	-	-	(SMD -0.59; 95% CI -1.01 to -0.16)
03 Betametha- sone 17-valerate 21-acetate plus tretinoine plus salicylic acid	Effect size [CI]	(SMD -0.76; 95% CI -1.21 to -0.31)	-	(SMD -0.54; 95% CI -0.99 to -0.10)	· · · · ·	(SMD -0.76; 95% CI -1.21 to -0.31)
04 Caffeine (top- ical) 10%, TD	Effect size [CI]	-	-	(SMD -0.39; 95% CI -0.84 to 0.06)	-	(SMD -0.39; 95% CI -0.84 to 0.06)
05 Calcipotriene 0. 005% ointment + nicotinamide 0.05% or 0.1%	Effect size [CI]	-	(SMD -0.48; 95% CI -0.81 to -0.15)	-	-	(SMD -0.48; 95% CI -0.81 to -0.15)

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Table 9.	Analysis 06: Tria	l characteristics and	outcomes: other treatments v	s. placebo	(Continued)
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or 0.7% or 1. 4%, BD						
06 Dead sea salts emollient lotion	Effect size [CI]	-	-	(SMD 0.57; 95% CI -0.36 to 1.51)	-	(SMD 0.57; 95% CI -0.36 to 1.51)
07 Fish oil plus occlusion	Effect size [CI]	-	(SMD -1.05; 95% CI -1.64 to -0.46)	-	-	(SMD -1.05; 95% CI -1.64 to -0.46)
08 Herbal skin care (Dr Michaels® cleans- ing gel, ointment and skin condi- tioner), BD	Effect size [CI]			(SMD -2.96; 95% CI -4.19 to -1.74)	-	(SMD -2.96; 95% CI -4.19 to -1.74)
09 Hexafluoro- 1,25-dihydrox- yvitamin D3	Effect size [CI]	(SMD -0.62; 95% CI -1.35 to 0.12)	(SMD -1.13; 95% CI -1.91 to -0.35)	-	-	(SMD -0.62; 95% CI -1.35 to 0.12)
10 Indigo natu- ralis 1.4% oint- ment	Effect size [CI]	(SMD -2.14; 95% CI -2.74 to -1.53)	(SMD -1.64; 95% CI -2.13 to -1.15)	-	-	(SMD -2.09; 95% CI -2.62 to -1.56)
11 Kukui nut oil, TD	Effect size [CI]		•	(SMD -0.03; 95% CI -0.84 to 0.77)	(SMD 0.00; 95% CI -0.80 to 0.80)	(SMD 0.00; 95% CI -0.80 to 0.80)
12 Ma- honia aquifolium (Reliéva™), BD	Effect size [CI]	-	-	-	-	(SMD -0.77; 95% CI -1.06 to -0.48)
13 Methotrexate gel	Effect size [CI]		(SMD -0.48; 95% CI -0.92 to -0.04)	(SMD -1.58; 95% CI -2.16 to -0.99)	-	(SMD -1.05; 95% CI -2.04 to -0.06)
14 Mycopheno- lic acid ointment	Effect size [CI]	-	(SMD -1.44; 95% CI -2.67 to -0.22)	-	-	(SMD -1.44; 95% CI -2.67 to -0.22)
15 NG- monomethyl-L- arginine (L- NMMA) cream	Effect size [CI]	-	(SMD 0.08; 95% CI -0.60 to 0.75)	-	-	(SMD 0.08; 95% CI -0.60 to 0.75)

Table 9.	Analysis 06: Tria	l characteristics and	l outcomes: other	treatments vs.	placebo	(Continued)
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16 Nicotinamide 1. 4%, BD	Effect size [CI]	-	(SMD -0.20; 95% CI -0.60 to 0.20)	-	-	(SMD -0.20; 95% CI -0.60 to 0.20)
17 Oleum hor- wathiensis	Effect size [CI]	(SMD -0.02; 95% CI -0.63 to 0.58)	(SMD -0.77; 95% CI -1.40 to -0.14)	-	-	(SMD -0.02; 95% CI -0.63 to 0.58)
18 Omega-3- polyunsat- urated fatty acids ointment	Effect size [CI]	-	-	-	-	-
19 Platelet aggre- gation activating factor (PAF) (Ro 24-0238)	Effect size [CI]	(SMD -0.07; 95% CI -0.50 to 0.37)	-		-	(SMD -0.07; 95% CI -0.50 to 0.37)
20 Polymyxin B cream, 200,000 U/g	Effect size [CI]	-	(SMD 0.13; 95% CI -0.59 to 0.85)	-	-	(SMD 0.13; 95% CI -0.59 to 0.85)
21 PTH (1-34) in Nova- some A® liposo- mal cream, BD	Effect size [CI]	-	(SMD -2.31; 95% CI -3.26 to -1.36)	-	-	(SMD -2.31; 95% CI -3.26 to -1.36)
22 Sirolimus (topi- cal), 2.2% for 6 wks, then 8% for a further 6 wks	Effect size [CI]	-	(SMD -0.39; 95% CI -0.98 to 0.21)	-	-	(SMD -0.39; 95% CI -0.98 to 0.21)
23 Tacrolimus ointment	Effect size [CI]	-	(SMD 0.06; 95% CI -0.52 to 0.63)	-	-	(SMD 0.06; 95% CI -0.52 to 0.63)
24 Tar	Effect size [CI]	-	(SMD -0.45; 95% CI -1.11 to 0.22)	-	-	(SMD -0.45; 95% CI -1.11 to 0.22)
25 Tazarotene	Effect size [CI]	-	(SMD -0.86; 95% CI -1.11 to -0.62)	-	-	(SMD -0.86; 95% CI -1.11 to -0.62)
26 Theophylline 1% ointment, BD	Effect size [CI]	-	-	(SMD -2.87; 95% CI -4.13 to -1.62)	-	(SMD -2.87; 95% CI -4.13 to -1.62)

All treatments	(not pooled)	-	-	-	-	-
-	No. participants	364	907	529	105	1450
-	Between-patient design	4	5	8	2	12
-	Within-patient design	4	12	1	0	14
-	Treatment dura- tion	3 wks to 12 wks	3 wks to 12 wks	2 wks to 12 wks	3 wks to 12 wks	2 wks to 12 wks

 Table 9. Analysis 06: Trial characteristics and outcomes: other treatments vs. placebo
 (Continued)

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Cal- cipotriol vs. be- tamethasone dipropionate	Effect size [CI]	(SMD 0.43; 95% CI 0. 28 to 0.58)	-	(SMD 0.36; 95% CI 0. 22 to 0.51)	-	(SMD 0.43; 95% CI 0. 28 to 0.58)
02 Calcipotriol vs. betametha- sone valerate	Effect size [CI]	(SMD -0.02; 95% CI -0.21 to 0.17)	(SMD -0.26; 95% CI -0.41 to -0.11)	(SMD -0.12; 95% CI -0.22 to -0.02)	(SMD -0.26; 95% CI -0.38 to -0.14)	(SMD -0.12; 95% CI -0.26 to 0.02)
03 Calcipotriol vs. desoxymeta- sone	Effect size [CI]	-	-	(SMD 0.15; 95% CI -0.73 to 1.02)	-	(SMD 0.15; 95% CI -0.73 to 1.02)
04 Calcipotriol vs. diflorasone diac- etate	Effect size [CI]	(SMD 0.27; 95% CI 0. 02 to 0.52)	(SMD 0.40; 95% CI 0. 15 to 0.65)	-	-	(SMD 0.27; 95% CI 0. 02 to 0.52)
05 Calcipotriol vs. fluocinonide	Effect size [CI]	(SMD -0.58; 95% CI -0.99 to -0.18)	(SMD -0.50; 95% CI -0.92 to -0.07)	-	-	(SMD -0.58; 95% CI -0.99 to -0.18)
06 Calcitriol vs. betamethasone dipropionate	Effect size [CI]	(SMD 0.21; 95% CI -0.04 to 0.45)	(SMD 0.27; 95% CI 0. 02 to 0.51)	(SMD 0.39; 95% CI 0. 14 to 0.63)	-	(SMD 0.21; 95% CI -0.04 to 0.45)

07 Calcitriol vs.	Effect size [CI]	(SMD -0.19;	-	-	-	(SMD -0.19;
betamethasone valerate		95% CI -0.91 to 0.53)				95% CI -0.91 to 0.53)
08 Tacalcitol vs. betamethasone valerate	Effect size [CI]	-	(SMD 0.41; 95% CI 0. 09 to 0.74)	-	-	(SMD 0.41; 95% CI 0. 09 to 0.74)
All treatments	Effect size [CI]; I ² statistic	95% CI -0.04 to	95% CI -0.22 to	(SMD 0.12; 95% CI -0.07 to 0.32); I ² statistic: 86.2%	(SMD -0.26; 95% CI - 0.38 to -0.14); I ² statistic: 0%	
-	No. participants	2655	891	3185	738	3542
-	Between-patient design	7	2	7	1	9
-	Within-patient design	1	4	2	1	5
-	Treatment dura- tion	3 wks to 8 wks	3 wks to 6 wks	4 wks to 8 wks	6 wks	3 wks to 8 wks
Sensitivity analy- ses	Within-patient trials	-	-	-	-	(SMD 0.17; 95% CI -0.20 to 0.54)
-	Between-patient trials	-	-	-	-	(SMD 0.10; 95% CI -0.11 to 0.31)
-	correlation coef- ficient (rho) = 0 All trials	-	-	-	-	(SMD 0.10; 95% CI -0.08 to 0.28); I ² statistic: 90. 5%
-	rho = 0 Btw-patient tri- als rho = 0.25 Within-patient trials	-	-	-	-	(SMD 0.10; 95% CI -0.07 to 0.28) I ² statistic: 91. 3%
-	rho = 0 Btw-patient tri- als rho = 0.50	-	-	-	-	(SMD 0.11; 95% CI -0.07 to 0.29)

Table 10. Analysis 07: Trial characteristics and outcomes: vitamin D vs. potent steroids (Continued)

Table 10. Analysis 07: Trial characteristics and outcomes: vitamin D vs. potent steroids (Continued)

Within-patient trials					I ² statistic: 92. 4%
- rho = 0 Btw-patient tri- als rho = 0.75 Within-patient trials	-	-	-	-	(SMD 0.12; 95% CI -0.06 to 0.30) I ² statistic: 94. 3%

Table 11.	Analysis 08: Trial characteristics and	l outcomes: vitamin D vs. v. potent steroids
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Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Calcipotriol vs. clobetasol propi- onate	Effect size [CI]	(SMD 0.19; 95% CI -0.42 to 0.80)	-	(SMD -0.32; 95% CI -0.95 to 0.30)	(SMD 0.42; 95% CI -0.20 to 1.03)	(SMD -0.06; 95% CI -0.57 to 0.44); I ² statistic: 25.7%
All treatments	No. participants	42	0	40	42	82
-	Between-patient design	1	0	1	1	2
-	Within-patient design	0	0	0	0	0
-	Treatment dura- tion	2 wks	-	6 wks	2 wks	2 wks to 6 wks

Table 12. Analysis 09: Trial characteristics and outcomes: vitamin D + steroid vs. steroid

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Calcipotriol + betamethasone dipropionate vs. betamethasone dipropionate	Effect size [CI]	(SMD -0.40; 95% CI -0.52 to -0.27)	-	(SMD -0.44; 95% CI -0.55 to -0.33)	-	(SMD -0.40; 95% CI -0.52 to -0.27)
02 Calcipotriol + betamethasone dipropionate vs.	Effect size [CI]	-	(SMD 0.45; 95% CI 0. 09 to 0.81)	-	-	(SMD 0.45; 95% CI 0.

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clobetasol propi- onate						09 to 0.81)
03 Calcipotriol + clobetasol propi- onate vs. clobeta- sol propionate	Effect size [CI]	(SMD -0.69; 95% CI -1.22 to -0.15)	-	-		(SMD -0.69; 95% CI -1.22 to -0.15)
-	Effect size [CI], I ²	-0.41; 95% CI -	95% CI 0.09 to	(SMD -0.44; 95% CI - 0.55 to -0.33) I ² statistic: 22.4%	95% CI -0.80 to	
-	No. participants	1991	122	1876	65	2113
-	Between-patient design	4	1	3	1	5
-	Within-patient design	0	0	0	0	0
-	Treatment dura- tion	2 wks to 8 wks	4 wks	4 wks to 8 wks	2 wks	2 wks to 8 wks

Table 12. Analysis 09: Trial characteristics and outcomes: vitamin D + steroid vs. steroid (Continued)

Table 13.	Analysis 10: 7	Frial characteristics	and outcomes:	vitamin D	vs. dithranol
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Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Calcipotriol vs. dithranol	Effect size [CI]		(SMD -0.54; 95% CI -1.16 to 0.08)	(SMD 0.73; 95% CI -0.55 to 2.00)		
02 Calcitriol vs. dithranol	Effect size [CI]	(SMD 0.51; 95% CI 0. 13 to 0.88)	•	(SMD -0.21; 95% CI -0.58 to 0.16)	-	(SMD 0.51; 95% CI 0. 13 to 0.88)
03 Tacalcitol vs. dithranol	Effect size [CI]	-		(SMD -0.07; 95% CI -0.50 to 0.36)	-	(SMD -0.18; 95% CI -0.60 to 0.25)
All treatments	Effect size [CI], I ² statistic	(SMD -0.24; 95% CI - 0.72 to 0.25); I ² statistic:93.0%		(SMD 0.36; 95% CI -0.33 to 1.04); I ² statistic: 94.5%		
-	No. participants	1108	386	796	544	1284

-	Between-patient design	5	3	5	2	7
-	Within-patient design	0	1	0	0	1
-	Treatment dura- tion	8 wks to 12 wks	4 wks to 8 wks	8 wks to 12 wks	8 wks to 12 wks	4 wks to 12 wks

Table 13. Analysis 10: Trial characteristics and outcomes: vitamin D vs. dithranol (Continued)

Table 14. Analysis 11: Trial characteristics and outcomes: vitamin D vs. vitamin D

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Calcipotriol vs. calcitriol	Effect size [CI]	(SMD 0.00; 95% CI -0.25 to 0.25)	(SMD -0.32; 95% CI -0.57 to -0.07)	(SMD -1.11; 95% CI -2.22 to 0.01)	(SMD 0.04; 95% CI -0.21 to 0.29)	(SMD -0.41; 95% CI -1.46 to 0.64)
02 Calcipotriol vs. tacalcitol	Effect size [CI]	×	(SMD -0.45; 95% CI -0.68 to -0.22)	-	-	(SMD -0.47; 95% CI -0.73 to -0.21)
03 Calcipotriol vs. maxacalcitol	Effect size [CI]	(SMD 0.43; 95% CI -0.12 to 0.98)	(SMD 0.13; 95% CI -0.41 to 0.68)	-	-	(SMD 0.43; 95% CI -0.12 to 0.98)
All treatments	Effect size [CI], I ²	(SMD -0.06; 95% CI - 0.51 to 0.38); I ² statistic: 82.2%		(SMD -1.11; 95% CI - 2.22 to 0.01); I ² statistic: NA	(SMD 0.04; 95% CI -0.21 to 0.29); I ² statistic: NA	(SMD -0.17; 95% CI - 0.62 to 0.27); I ² statistic: 78.5%
-	No. participants	498	563	15	250	513
-	Between-patient design	2	2	1	1	3
-	Within-patient design	1	1	0	0	1
-	Treatment dura- tion	8 wks to 12 wks	8 wks to 12 wks	8 wks	12 wks	8 wks to 12 wks
Sensitivity analy- ses	TSS data from Ji 2008 used in combined end point: 01 Calcipotriol		-	-	-	(SMD -0.52; 95% CI -1.19 to 0.15; I ² statistic: 44.9%)

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Table 14.	Analysis 11: Trial characteristics and outcomes: vitamin D vs. vit	tamin D (Continued)
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	vs. calcitriol				
-	TSS data from Ji 2008 used in combined end point: all treatments	-	-	-	(SMD -0.28; 95% CI -0.66 to 0.10; I ² statistic: 70.6%)

Table 15. Analysis 12: Trial characteristics and outcomes: vitamin D vs. vitamin D + steroid

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Cal- cipotriol BD vs. calcipotriol OM, BMD ON	Effect size [CI]	(SMD 0.56; 95% CI 0. 23 to 0.88)	-	(SMD 0.46; 95% CI 0. 10 to 0.82)	-	(SMD 0.56; 95% CI 0. 23 to 0.88)
02 Cal- cipotriol OD vs. combined cal- cipotriol + BMD OD	Effect size [CI]	(SMD 0.66; 95% CI 0. 31 to 1.02)	-	(SMD 0.67; 95% CI 0. 23 to 1.11)	-	(SMD 0.66; 95% CI 0. 31 to 1.02)
03 Cal- cipotriol BD vs. combined cal- cipotriol + BMD OD	Effect size [CI]	(SMD 0.27; 95% CI 0. 06 to 0.48)	(SMD 0.25; 95% CI 0. 03 to 0.48)	(SMD 0.52; 95% CI 0. 38 to 0.67)	-	(SMD 0.43; 95% CI 0. 20 to 0.66)
04 Cal- cipotriol BD vs. combined cal- cipotriol + BMD BD	Effect size [CI]	(SMD 0.66; 95% CI 0. 40 to 0.93)	-	(SMD 0.64; 95% CI 0. 46 to 0.83)	-	(SMD 0.66; 95% CI 0. 40 to 0.93)
05 Cal- cipotriol BD vs. calcipotriol OM, BMV ON	Effect size [CI]	(SMD 0.27; 95% CI -0.19 to 0.74)	-	(SMD 0.43; 95% CI -0.07 to 0.93)	-	(SMD 0.27; 95% CI -0.19 to 0.74)
06 Cal- cipotriol BD vs. calcipotriol OM, clobetasone bu- tyrate ON	Effect size [CI]	(SMD 0.27; 95% CI 0. 05 to 0.48)	-	(SMD 0.17; 95% CI -0.04 to 0.38)	-	(SMD 0.27; 95% CI 0. 05 to 0.48)

07 Cal- cipotriol BD vs. calcipotriol BD + clobetasol pro- pionate BD	Effect size [CI]	(SMD 0.88; 95% CI 0. 34 to 1.42)	-	-	(SMD 0.70; 95% CI 0. 16 to 1.23)	(SMD 0.88; 95% CI 0. 34 to 1.42)
08 Cal- cipotriol BD vs. calcipotriol OM, diflucortolone valerate ON	Effect size [CI]	-	-	(SMD 0.08; 95% CI -0.29 to 0.44)	-	(SMD 0.08; 95% CI -0.29 to 0.44)
09 Cal- cipotriol OD vs. calcipotriol OM, fluocinonide acetonide ON	Effect size [CI]	-	-	(SMD 0.53; 95% CI -0.11 to 1.18)	-	(SMD 0.53; 95% CI -0.11 to 1.18)
10 Calcipotriol OD vs. combined calcipotriol + hydrocortisone OD	Effect size [CI]	(SMD 0.14; 95% CI -0.06 to 0.33)	-	(SMD 0.08; 95% CI -0.11 to 0.28)	-	(SMD 0.14; 95% CI -0.06 to 0.33)
11 calcitriol BD vs. diflu- cortolone valer- ate OM, calcitriol ON	Effect size [CI]	-	-	(SMD 0.24; 95% CI -0.09 to 0.57)	-	(SMD 0.24; 95% CI -0.09 to 0.57)
12 Tacalci- tol OD vs. com- bined cal- cipotriol + BMD OD	Effect size [CI]	(SMD 0.48; 95% CI 0. 26 to 0.70)	-	(SMD 0.47; 95% CI 0. 25 to 0.69)	(SMD 0.46; 95% CI 0. 24 to 0.68)	(SMD 0.48; 95% CI 0. 26 to 0.70)
All treatments	Effect size [CI], I ² statistic		(SMD 0.25; 95% CI 0. 03 to 0.48)		(SMD 0.49; 95% CI 0.29 to 0.69), I ² statistic: 0%	(SMD 0.46; 95% CI 0.33 to 0.59), I ² statistic: 83.3%
-	No. participants	4791	301	5703	399	5856
-	Between-patient design	11	1	15	2	16
-	Within-patient design	0	0	1	0	1

Table 15. Analysis 12: Trial characteristics and outcomes: vitamin D vs. vitamin D + steroid (Continued)

-	Treatment dura- tion	2 wks to 8 wks	4 wks	2 wks to 12 wks	2 wks to 8 wks	2 wks to 12 wks
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Table 15.	Analysis 12: Tria	l characteristics and	outcomes: vitamin	D vs. vitamin D	+ steroid	(Continued)
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Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Calcipotriol (12 wks) vs. com- bined cal- cipotriol + BMD (8 wks); then cal- cipotriol (4 wks)	Effect size [CI]	(SMD -0.12; 95% CI -0.29 to 0.04)	-	(SMD -0.04; 95% CI -0.19 to 0.11)	(SMD -0.14; 95% CI -0.30 to 0.02)	(SMD -0.12; 95% CI -0.29 to 0.04)
02 Calcipotriol (12 wks) vs. com- bined cal- cipotriol + BMD (4 wks); then cal- cipotriol (8 wks)	Effect size [CI]	-	-	(SMD 0.29; 95% CI -0.04 to 0.62)	-	(SMD 0.29; 95% CI -0.04 to 0.62)
03 Calcipotriol (12 wks) vs. com- bined cal- cipotriol + BMD (4 wks); then cal- cipotriol (w/dy) & combined cal- cipotriol + BMD (w/e) (8 wks)	Effect size [CI]	(SMD 0.13; 95% CI -0.04 to 0.29)	-	(SMD 0.10; 95% CI -0.05 to 0.25)	(SMD 0.10; 95% CI -0.06 to 0.26)	(SMD 0.13; 95% CI -0.04 to 0.29)
04 Calcipotriol (6 wks) vs. clobe- tasol propionate (2 wks); then cal- cipotriol (4 wks)	Effect size [CI]	(SMD 0.60; 95% CI 0. 18 to 1.02)	(SMD 0.63; 95% CI 0. 21 to 1.05)	-	-	(SMD 0.60; 95% CI 0. 18 to 1.02)
05 Calcipotriol (6 wks) vs. calcipotriol OM, fluocinonide	Effect size [CI]	-	-	(SMD 0.66; 95% CI 0. 01 to 1.32)	-	(SMD 0.66; 95% CI 0. 01 to 1.32)

Table 16.	Analysis 13: Tria	l characteristics and	l outcomes: vitamin D	vs. other treatments:	complex regimens
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acetonide ON (2 wks); then cal- cipotriol BD (4 wks)						
06 Cal- cipotriol (6 wks) vs. halometasone OM, calcipotriol ON (2 wks); then cal- cipotriol BD (w/ dy), halometa- sone (w/ e) (2 wks); then calcipotriol BD (2wks)	Effect size [CI]	(SMD 0.41; 95% CI -0.05 to 0.86)	-	(SMD 1.13; 95% CI 0. 64 to 1.62)	-	(SMD 0.41; 95% CI -0.05 to 0.86)
07 Calcipotriol ON, clobetasol propionate OM (2 to 4 wks); then calcipotriol BD (to wk12) vs. calcitriol ON, clobetasol propi- onate OM (2 to 4 wks); then cal- citriol BD (to wk12)	Effect size [CI]	(SMD -0.19; 95% CI -0.54 to 0.16)	-	(SMD -0.27; 95% CI -0.62 to 0.09)	-	(SMD -0.19; 95% CI -0.54 to 0.16)
08 Combined calcipotriol + BMD (4 wks) ; then placebo ointment BD (8 wks) vs. com- bined cal- cipotriol + BMD (4 wks); then calcipotriol oint- ment BD (8 wks)	Effect size [CI]	(SMD 0.27; 95% CI 0. 12 to 0.41)	-	(SMD 0.25; 95% CI 0. 10 to 0.39)	(SMD 0.28; 95% CI 0. 13 to 0.42)	(SMD 0.27; 95% CI 0. 12 to 0.41)
09 Combined calcipotriol + BMD (4 wks) ; then placebo ointment BD (8 wks) vs. com- bined cal-	Effect size [CI]	(SMD 0.51; 95% CI 0. 37 to 0.66)	-	(SMD 0.59; 95% CI 0. 45 to 0.74)	(SMD 0.71; 95% CI 0. 56 to 0.85)	(SMD 0.51; 95% CI 0. 37 to 0.66)

Table 16. Analysis 13: Trial characteristics and outcomes: vitamin D vs. other treatments: complex regimens (Continued)

cipotriol + BMD (4 wks); then cal- cipotriol (w/dy) + combined cal- cipotriol + BMD (w/e) (8 wks)						
10 combined calcipotriol + BMD (4 wks); then calcipotriol ointment BD (8 wks) vs. com- bined cal- cipotriol + BMD (4 wks); then cal- cipotriol (w/dy) + combined cal- cipotriol + BMD (w/e) (8 wks)	Effect size [CI]	(SMD 0.26; 95% CI 0. 11 to 0.40)	-	(SMD 0.30; 95% CI 0. 16 to 0.45)	(SMD 0.44; 95% CI 0. 29 to 0.58)	(SMD 0.26; 95% CI 0. 11 to 0.40)
11 Combined calcipotriol + BMD (8 wks); then calcipotriol (4 wks) vs. com- bined cal- cipotriol + BMD (4 wks); then cal- cipotriol (w/dy) & combined cal- cipotriol + BMD (w/e) (8 wks)	Effect size [CI]	(SMD 0.24; 95% CI 0. 08 to 0.40)	-	(SMD 0.15; 95% CI -0.01 to 0.30)	(SMD 0.23; 95% CI 0. 07 to 0.39)	(SMD 0.24; 95% CI 0. 08 to 0.40)
12 Tacalcitol (8 wks) vs. com- bined cal- cipotriol + BMD (4 wks); then cal- cipotriol (4 wks)	Effect size [CI]	(SMD 0.54; 95% CI 0. 36 to 0.72)	-	(SMD 0.49; 95% CI 0. 31 to 0.67)	(SMD 0.54; 95% CI 0. 36 to 0.72)	(SMD 0.54; 95% CI 0. 36 to 0.72)
All treatments	(not pooled)	-	-	-	-	-
-	No. participants	2755	46	2991	2508	2936
-	Between-patient design	6	0	8	4	8

Table 16. Analysis 13: Trial characteristics and outcomes: vitamin D vs. other treatments: complex regimens (Continued)

-	Within-patient design	1	1	0	0	1
-	Treatment dura- tion	6 wks to 12 wks	6 wks	2 wks to 12 wks	8 wks to 12 wks	2 wks to 12 wks
For acronyms, see	Table 1.					

Table 16. Analysis 13: Trial characteristics and outcomes: vitamin D vs. other treatments: complex regimens (Continued)

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Table 17. Analysis 14: Trial characteristics and outcomes: vitamin D vs. other treatment: long-term studies (> 24 wks)

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Combined cal- cipotriol + BMD (52 wks) vs. al- ternating: com- bined cal- cipotriol + BMD (4 wks); then cal- cipotriol (4 wks)	Effect size [CI]	(SMD -0.09; 95% CI -0.36 to 0.18)	-	-	-	(SMD -0.09; 95% CI -0.36 to 0. 18)
02 Combined cal- cipotriol+BMD (52 wks) vs. combined cal- cipotriol+ BMD (4 wks); then calcipotriol (48 wks)	Effect size [CI]	(SMD -0.18; 95% CI -0.47 to 0.10)	-	-	-	(SMD -0.18; 95% CI -0.47 to 0. 10)
03 Alternating: combined cal- cipotriol + BMD (4 wks); then cal- cipotriol (4 wks) vs. combined calcipotriol + BMD (4 wks); then calcipotriol (48 wks)	Effect size [CI]	(SMD -0.09; 95% CI -0.37 to 0.19)	-	-	-	(SMD -0.09; 95% CI -0.37 to 0. 19)
All treatments	(no pooling)	-	-	-	-	-
-	No. participants	297	0	0	0	297

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-	Between-patient design	1	0	0	0	1
-	Within-patient design	0	0	0	0	0
-	Treatment dura- tion	52 wks	-	-	-	52 wks

Table 17. Analysis 14: Trial characteristics and outcomes: vitamin D vs. other treatment: long-term studies (> 24 wks) (Continued)

Table 18. Analysis 15: Trial characteristics and outcomes: vitamin D/other treatment

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Calcipotriol vs. coal tar	Effect size [CI]	(SMD -0.53; 95% CI -1.74 to 0.68)	-	(SMD -0.10; 95% CI -1.54 to 1.35)	(SMD -0.10; 95% CI -1.54 to 1.35)	(SMD -0.53; 95% CI -1.74 to 0.68)
02 Calcipotriol vs. coal tar poly- therapy	Effect size [CI]	(SMD -0.59; 95% CI -0.87 to -0.31)	(SMD -0.51; 95% CI -0.86 to -0.16)	(SMD -0.63; 95% CI -1.06 to -0.20)	(SMD -0.63; 95% CI -1.06 to -0.20)	(SMD -0.59; 95% CI -0.87 to -0.31)
03 Calcipotriol vs. nicotinamide 1.4%, BD	Effect size [CI]	-	(SMD -0.09; 95% CI -0.49 to 0.31)	-	-	(SMD -0.09; 95% CI -0.49 to 0.31)
04 Calcipotriol vs. calcipotriol + nicotinamide 1. 4%, BD	Effect size [CI]	-	(SMD 0.19; 95% CI -0.14 to 0.52)	-	-	(SMD 0.19; 95% CI -0.14 to 0.52)
05 Calcipotriol vs. corticosteroid + salicylic acid	Effect size [CI]	(SMD -0.06; 95% CI -0.33 to 0.22)	-	(SMD -0.05; 95% CI -0.36 to 0.26)	(SMD -0.49; 95% CI -0.79 to -0.20)	(SMD -0.05; 95% CI -0.26 to 0.15)
06 Calcipotriol vs. propylth- iouracil cream	Effect size [CI]	-	-	(SMD -2.24; 95% CI -3.23 to -1.25)	-	(SMD -2.24; 95% CI -3.23 to -1.25)
07 Calcipotriol vs. tacrolimus oint- ment	Effect size [CI]	-	(SMD -0.35; 95% CI -1.51 to 0.81)			(SMD -0.55; 95% CI -1.28 to 0.17)

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Table 18.	Analysis 15: Tria	l characteristics and	l outcomes: vitamin D/ot	her treatment	(Continued)
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08 Calcipotriol vs. tazarotene	Effect size [CI]	(SMD -0.22; 95% CI -0.60 to 0.16)		-	(SMD -0.35; 95% CI -0.99 to 0.29)	(SMD -0.10; 95% CI -0.35 to 0.16)
09 Calcipotriol vs. tazarotene gel plus mometasone furoate cream	Effect size [CI]	-	-	-	-	-
10 Calcipotriol vs. vitamin B12 cream	Effect size [CI]	(SMD -0.55; 95% CI -1.33 to 0.24)	-	(SMD -0.01; 95% CI -0.78 to 0.75)		(SMD -0.55; 95% CI -1.33 to 0.24)
11 Head-to- head vitamin D alone or in com- bination: dosing	Effect size [CI]	(SMD -0.24; 95% CI -0.38 to -0.09)	-	(SMD -0.12; 95% CI -0.25 to 0.00)	-	(SMD -0.20; 95% CI -0.32 to -0.07)
12 Head-to- head vitamin D alone or in com- bination: occlu- sion	Effect size [CI]	-	(SMD -0.18; 95% CI -2.04 to 1.68)	-	-	(SMD -0.18; 95% CI -2.04 to 1.68)
All treatments	(not pooled)	-	-	-	-	-
-	No. participants	1386	898	1228	456	2364
-	Between-patient design	8	5	6	3	13
-	Within-patient design	2	2	3	3	6
-	Treatment dura- tion Table 1	4 wks to 12 wks	6 wks to 12 wks	4 wks to 12 wks	4 wks to 12 wks	4 wks to 12 wks

Table 19. Analysis 16: Trial characteristics and outcomes: flexural/facial psoriasis: placebo trials

01 Betametha- sone valerate 0. 1%, OD	Effect size [CI]	-	-	(SMD -2.83; 95% CI -3.79 to -1.88)	-	(SMD -2.83; 95% CI -3.79 to -1.88)
02 Calcipotriol ointment, OD	Effect size [CI]	-	-	(SMD -1.08; 95% CI -1.77 to -0.40)	-	(SMD -1.08; 95% CI -1.77 to -0.40)
03 Pimecrolimus cream, 1% OD/ BD	Effect size [CI]	(SMD -1.07; 95% CI -1.69 to -0.45)		(SMD -0.62; 95% CI -1.27 to 0.02)	(SMD -0.65; 95% CI -1.24 to -0.06)	(SMD -0.86; 95% CI -1.30 to -0.41)
04 Tacrolimus oint- ment 0.1%, BD	Effect size [CI]	-	-	-	-	-
All treatments	(no pooling)	-	-	-	-	-
-	No. participants	47	57	75	47	122
-	Between-patient design	1	1	1	1	2
-	Within-patient design	0	0	0	0	0
-	Treatment dura- tion	8 wks	8 wks	4 wks	8 wks	4 wks to 8 wks

 Table 19. Analysis 16: Trial characteristics and outcomes: flexural/facial psoriasis: placebo trials (Continued)

Table 20. Analysis 17: Trial characteristics and outcomes: flexural/facial psoriasis: vitamin D vs. other treatment

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Calcipotriol vs. BMV	Effect size [CI]	-	-	(SMD 2.02; 95% CI 1. 20 to 2.84)	-	(SMD 2.02; 95% CI 1.20 to 2.84)
02 Calcipotriol vs. calcipotriol + hydrocortisone	Effect size [CI]	(SMD 0.30; 95% CI 0. 11 to 0.50)	-	(SMD 0.32; 95% CI 0. 12 to 0.51)	-	(SMD 0.30; 95% CI 0.11 to 0.50)
03 Calcipotriol vs. calcitriol	Effect size [CI]	-	(SMD 0.61; 95% CI 0. 28 to 0.94)	-	-	(SMD 0.61; 95% CI 0.28 to 0.94)

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04 Calcipotriol vs. pimecrolimus	Effect size [CI]	-	-	(SMD -0.53; 95% CI -1.17 to 0.11)		(SMD -0. 53; 95% CI -1.17 to 0.11)
05 Calcitriol vs. tacrolimus	Effect size [CI]		(SMD 0.29; 95% CI -0.27 to 0.85)	-	-	(SMD 0.42; 95% CI -0.15 to 0.98)
All treatments	(no pooling)	-	-	-	-	-
-	No. participants	457	124	464	0	588
-	Between-patient design	2	1	2	0	3
-	Within-patient design	0	1	0	0	1
-	Treatment dura- tion	6 wks to 8 wks	6 wks	4 wks to 8 wks	0	4 wks to 8 wks

 Table 20. Analysis 17: Trial characteristics and outcomes: flexural/facial psoriasis: vitamin D vs. other treatment (Continued)

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Vitamin D: calcipotriol	Effect size [CI]	(SMD -0.72; 95% CI -1.28 to -0.16)	95% CI -0.64 to		(SMD -0.66; 95% CI -1.28 to -0.05)	
02 Po- tent steroid: be- tamethasone dipropionate	Effect size [CI]	(SMD -1.09; 95% CI -1.29 to -0.90)	95% CI -1.19 to		(SMD -1.23; 95% CI -1.43 to -1.03)	
03 Po- tent steroid: be- tamethasone valerate	Effect size [CI]	-	(SMD -1.40; 95% CI -1.75 to -1.05)	-	-	(SMD -1.40; 95% CI -1.75 to -1.05)
04 Very po- tent steroid: am- cinonide	Effect size [CI]		(SMD -1.58; 95% CI -1.98 to -1.18)		(SMD -0.97; 95% CI -1.33 to -0.61)	

Table 21. Analysis 18: Trial characteristics and outcomes: scalp psoriasis: placebo-controlled tria	Table 21.
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05 Very potent steroid: clobeta- sol propionate	Effect size [CI]	(SMD -1.73; 95% CI -1.99 to -1.48)	•	-	-	(SMD -1.57; 95% CI -1.81 to -1.34)
06 Very po- tent steroid: hal- cinonide	Effect size [CI]	(SMD -1.11; 95% CI -1.69 to -0.53)	-	-	-	(SMD -1.11; 95% CI -1.69 to -0.53)
07 Vitamin D in combina- tion: calcipotriol + BMD	Effect size [CI]	(SMD -0.97; 95% CI -1.61 to -0.32)	(SMD -0.92; 95% CI -1.42 to -0.43)	-	(SMD -1.00; 95% CI -1.79 to -0.22)	(SMD -0.97; 95% CI -1.61 to -0.32)
08 Other treat- ment: be- tamethasone-17, 21-dipropionate plus salicylic acid	Effect size [CI]	(SMD -1.48; 95% CI -2.50 to -0.47)	(SMD -1.15; 95% CI -2.11 to -0.19)	-	-	(SMD -1.48; 95% CI -2.50 to -0.47)
09 Other treat- ment: ci- clopirox olamine shampoo	Effect size [CI]	-	(SMD -0.07; 95% CI -0.82 to 0.68)	-	(SMD -0.11; 95% CI -0.86 to 0.64)	(SMD -0.07; 95% CI -0.82 to 0.68)
10 Other treat- ment: fluoci- nolone ace- tonide, plus oc- clusion	Effect size [CI]	(SMD -1.22; 95% CI -1.69 to -0.76)	(SMD -0.89; 95% CI -1.34 to -0.44)	-	-	(SMD -1.22; 95% CI -1.69 to -0.76)
11 Other treat- ment: salicylic acid	Effect size [CI]	(SMD -0.86; 95% CI -1.79 to 0.06)	(SMD -0.57; 95% CI -1.47 to 0.32)	-	-	(SMD -0.86; 95% CI -1.79 to 0.06)
All treatments	No. participants	2472	2897	0	1875	3011
-	Between-patient design	9	12	0	5	13
-	Within-patient design	1	0	0	0	1
-	Treatment dura- tion	2 wks to 8 wks	2 wks to 8 wks	-	3 wks to 8 wks	2 wks to 8 wks
Sensitivity analy- sis: potent corti- costeroids	Effect size [CI]; I ² statistic	-	-	-	-	(SMD -1.18; 95% CI -1.40 to -0.96); I ² statistic: 19.9%

Table 21. Analysis 18: Trial characteristics and outcomes: scalp psoriasis: placebo-controlled trials (Continued)

Table 21. Analysis 18: Trial characteristics and outcomes: scalp psoriasis: placebo-controlled trials (Continued)

For acronyms, see Table 1.

Table 22. Analysis 19: Trial characteristics and outcomes: scalp psoriasis: vitamin D vs. other treatment	Table 22.	Analysis 19: Trial characteristics and outcomes: scalp psoriasis: vitamin D vs. other treatment
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Subcategory	Measure 01 IAGI/IGA 02 TSS 03 PASI 04 PA		04 PAGI/PGA	05 Combined end point		
01 Vitamin D vs. corticosteroid (potent) : calcipotriol vs. BMD	Effect size [CI]	size [CI] (SMD (SMD - (SMD 0.56; 95% CI 0. 0.48; 95% CI 0. 32 to 0.64) 28 to 0.63) - (SMD 0.56; 95% CI 0. 28 to 0.63)		(SMD 0.48; 95% CI 0.32 to 0.64)		
02 Vitamin D vs. corticosteroid (potent) : calcipotriol vs. BMV	Effect size [CI]			(SMD 0.37; 95% CI 0.20 to 0.55)		
03 Vitamin D vs. corticosteroid (very potent) : calcipotriol vs. clobetasol propi- onate	Effect size [CI]	-	(SMD 0.37; 95% CI 0. 05 to 0.69)	-	-	(SMD 0.37; 95% CI 0.05 to 0.69)
04 Vitamin D + corticosteroid vs. cor- ticosteroid: cal- cipotriol + BMD vs. BMD	porticosteroid vs. 9 por4 costeroid: cal- potriol + BMD		(SMD -0.19; 95% CI -0.27 to -0.11)	-	(SMD -0.17; 95% CI -0.25 to -0.09)	(SMD -0.18; 95% CI -0.26 to -0.10)
05 Vitamin D vs. vitamin D + cor- ticosteroid: cal- cipotriol vs. cal- cipotriol + BMD	Effect size [CI]	(SMD 0.64; 95% CI 0. 44 to 0.84)	(SMD 0.70; 95% CI 0. 56 to 0.84)	-	(SMD 0.84; 95% CI 0.61 to 1.08)	(SMD 0.64; 95% CI 0.44 to 0.84)
06 Vitamin D vs. other treat- ments: cal- cipotriol vs. coal tar polytherapy	Effect size [CI]	(SMD -0.24; 95% CI -0.73 to 0.25)	(SMD -0.30; 95% CI -0.84 to 0.24)	-	-	(SMD -0.45; 95% CI -0.92 to 0.02)

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All treatments	No. participants	5175	4877	0	3742	5413
-	Between-patient design	10	11	0	6	12
-	Within-patient design	0	0	0	0	0
-	Treatment dura- tion	4 wks to 52 wks	4 wks to 8 wks	0	4 wks to 8 wks	4 wks to 52 wks

Table 22. Analysis 19: Trial characteristics and outcomes: scalp psoriasis: vitamin D vs. other treatment (Continued)

Table 23. Analysis 04: Trial characteristics and outcomes: dithranol vs. placebo

Subcategory	Measure	01 IAGI/IGA	02 TSS	03 PASI	04 PAGI/PGA	05 Combined end point
01 Dithranol	Effect size [CI], N, I ² statistic	-	(SMD -1. 06; 95% CI -1.66 to -0.46); I ² statis- tic: 37.4%	-	-	(SMD -1.06; 95% CI -1.66 to -0.46); I ² statistic: 37.4%
-	No. participants	0	47	0	0	47
-	Between-patient design	0	0	0	0	0
-	Within-patient design	0	3	0	0	3
-	Treatment dura- tion		3 wks to 8 wks	-	-	3 wks to 8 wks
-	correlation coef- ficient (rho) = 0 All trials	-		-	-	(SMD -0.98; 95% CI -1.56 to -0.41) I ² statistic: 13.9%
-	rho = 0 Btw-patient trials rho = 0.25 Within-patient trials	-	-	-	-	(SMD -1.05; 95% CI -1.67 to -0.44) I ² statistic: 35.4%
-	rho = 0 Btw-patient trials rho = 0.50 Within-patient	-	-	-	-	(SMD -1.12; 95% CI -1.75 to -0.48) I ² statistic: 56.9%

Topical treatments for chronic plaque psoriasis (Review)

 Table 23. Analysis 04: Trial characteristics and outcomes: dithranol vs. placebo
 (Continued)

trials					
- rho = 0 Btw-patient trials rho = 0.75 Within-patient trials	-	-	-	-	(SMD -1.17; 95% CI -1.81 to -0.52) I ² statistic: 78.5%

Table 24. Included studies of adverse events

Study	Methods	Participants	Intervention (s)	Outcomes (AEs)	Summary findings	Notes	Allocation concealment
Andres 2006	DESIGN: be- tween-patient patient deliv- ery ALLOCA- TION: random Method of randomisa- tion: computer- generated list Concealment: unclear BLIND- ING: single- blind (investi- gator) WITH- DRAWAL/ DROPOUT: described	TD: 4 wks; FU: 4 wks LF: 0 (0%) BC: character- istics reported, but not demonstrated to be compa- rable (sham- poo group had more severe disease and higher proportion of	15 minutes Clobetasol propionate 0.	trasound mea- surement of skin thickness (epidermis + dermis) (mm) , averaged over 3 sites of the scalp) ocular sa- fety (intraocu- lar pressure) DSS (10- pt; sum of ery- thema, adher- ent desquama- tion, and	ther formula- tion had an impact on oc- ular safety, no report of ocu- lar stinging LAE: CS: 1/14; CG: 2/14 HPA suppres- sion: CS: 0/14; CG: 2/14 Atrophy: CS: 0/14; CG: 0/14 De- crease in skin thickness from baseline: mean difference: CS: 0.04 mm CG: -0.24 mm (difference: P < = 0.025)	Galderma	Unclear

		DSS > = 3 EXCLU- SION CRI- TERIA: preg- nancy or risk thereof; lacta- tion; ophthal- mological dis- order; contact lens wearer			similar. Com- pli- ance with pro- tocol was good in both groups		
Barnes 2000	DESIGN: within-patient patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	FU: 52 wks LF: 64 (32%) BC: NA Age: 46 (14. 5SD) Gender (per cent men) : 60%	Cal- cipotriol scalp solution 50 mcg/ml BD plus calcipotriol cream 50 mcg/ g BD (up to 70 g/wk) No control	Local AEs: number of AEs/partic- ipant % severe AEs withdrawals due to adverse events (WA) Systemic AEs: serum calcium serum PTH uri- nary calcium/ creatinine ra- tio	mon local AE was facial irri- tation (60/202 par- ticipants at wk 2), though the incidence de- clined rapidly over time (1/ 141 at wk 46). 20% of local AEs consid-	Leo Pharma-	Not applicable

Table 24. Included studies of adverse events	(Continued)
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		EXCLU- SION CRI- TERIA: preg- nancy or risk thereof; severe (i.e. requiring ad- ditional ther- apy) or un- stable psoria- sis; hyper- calcaemia; his- tory of hypo- or hy- perparathy- roidism, renal/ hepatic dis- ease; systemic or photothera- pies within previous 6 wks; other medication that could af- fect course of disease					
Berth-Jones 1993; Berth- Jones 1992c	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: NA	LF: 0 (0%) BC: NA Age: 43 Gender (per cent men) : 65% Severity: mean	Study A: cal- cipotriol oint- ment 50 mcg/ g BD up to 100 g/wk No control Study B: cal- cipotriol oint- ment 50 mcg/ g BD, using 100 g/wk for 4 wks Control: peo- ple using al- ternative ther- apies	not assessed Systemic AEs: urine calcium and phosphate ex- cretion; serum total calcium, phosphate and alkaline phos-	no significant trend in urine calcium excre- tion Study B: significant in- crease in urine calcium excre-	For study B, baseline com- pa- rability of in-	Not applicable

		demonstrated Age: 48 {42} Gender (per cent men) : 50% {44%} Severity: mean PASI: 18.0 (13.9SD) {NR} INCLUSION CRITERIA: people with chronic plaque psoria- sis; aged \geq 18; un- der long-term care of inves- tigators. Study A: compliant patients, responsive to calcipotriol. Study B: more extensive disease, failing to respond to low doses of topical agents. Controls re- ceived no cal- cipotriol. EXCLU- SION CRI- TERIA: preg- nancy				
Bleiker 1998	DESIGN: un- controlled study Delivery: un- clear ALLOCA- TION: non- random Method of randomisa- tion: NA	TD: 2 wks; FU: 26 wks	STUDY A: 200 g cal- cipotriol oint- ment 50 mcg/ g (wk 1) plus 300 g 50 mcg/ g calcipotriol (wk 2) STUDY B: Calcipotriol	Local AEs: not assessed Systemic AEs: serum total adjusted calcium urinary calcium	5 participants developed hy- percal- caemia during treatment, all had received a dose > 5 g/kg 9 participants became hyper- calciuric dur- ing treatment;	 Not applicable

	Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: not described	Sever- ity: PASI: 21.4 (range: 8.2 to 53.7) INCLUSION CRI- TERIA: inpa- tients with se- vere chronic plaque psori- asis (> 15% BSA) EXCLU- SION CRI- TERIA: renal impairment, pregnancy, lacta- tion, systemic treatment, di- uretics			this was un- correlated with dose		
Brodell 2011b	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non- randomised BLINDING: open WITH- DRAWAL/ DROPOUT: not described	N: 305 TD: 12 wks; FU: 12 LF: unclear BC: NA Age: 50.3 (13. 7SD); range: 22 to 84 Gender (per cent men): 61. 8% Severity: ODS: all pa- tients scored as moderate/ severe/very se- vere % BSA: 7.1% % white: 91. 8% INCLUSION CRI- TERIA: peo- ple with mod- erate to severe plaque psori- asis; affected;	Clo- betasol propi- onate 0.05% spray BD (2 to 4 wks); treat- ment respon- ders (ODS < = 3) then treated with calcitriol 3 mg/g oint- ment (8 wks)	tasias, burn- ing/stinging (0 to 3), skin atrophy, folli- culitis Overall disease sever- ity (ODS) (5- pt: 0 = clear to 4 = severe/very	telangiectasias 2/285 stinging/ Burning 39/ 285 folliculitis 11/ 285 At 12 wks: skin atrophy	Sponsored by Galderma lab- oratories	Not applicable

		aged 18 to 80 EXCLU- SION CRITERIA: not stated					
Corbett 1976	DESIGN: within-patient patient deliv- ery ALLOCA- TION: random Method of randomisa- tion: NR Concealment: unclear BLIND- ING: double- blind (partici- pant/ investigator) WITH- DRAWAL/ DROPOUT: not described	LF: 2 (14.3%) BC (clinical): NR Age: 44 (18. 4SD) Gender (per cent men) : 64% Severity: NR INCLUSION CRITERIA: bilateral psori-	Clobeta- sol propionate 0.05% oint- ment, BD Be- tamethasone valerate 0.1% ointment, BD	NR Systemic AEs: synacthen test for function of pituitary-	Quantities used by study par- ticipants were small (mean: 7 g/wk) No pituitary- adrenal suppression observed	Sponsorship not reported	Unclear
Gerritsen 2001; Langner 1996; van de Kerkhof 1996c	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: NA Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	TD: < = 78 wks; FU: <=	Calcitriol 3 mcg/g BD No control	Local AEs: serious AEs re- ported; with- drawals due to adverse events (WA) Systemic AEs: laboratory lev- els for: pro- tein albumin; calcium, phos- phorus, sodium, potas- sium, plasma calcitriol Urinary calcium, cre-	no se- rious local ad- verse events Systemic AEs: WA: 1/253. 4 addi- tional partic- ipants experi- enced hyper- calcaemia. All mean values for all parameters re- mained within	Sponsored by Solvay- Duphar BV Excludes scalp	Not applicable

Table 24.	Included	studies	of adverse events	(Continued)
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		sis; aged ≥18 EXCLU- SION CRI- TERIA: non- compliant; preg- nancy; use of systemic/ phototherapy within previ- ous 2 mths; use of topical therapy within previ- ous 1 wk; con- comitant dis- ease; clinically rel- evant abnor- mality in lab- oratory assess- ments; known hypersen- sitivity to vita- min D/vehicle		atinine, phos- phorus; creati- nine clearance; uri- nary calcium/ creatinine ra- tio	Mean use: 6 g/ day (range: 1 to 24 g/day)		
Guzzo 1996	DESIGN: be- tween-patient patient deliv- ery ALLOCA- TION: random Method of randomisa- tion: NR Concealment: unclear BLIND- ING: double- blind (partici- pant/ investigator) WITH- DRAWAL/ DROPOUT: described	TD: 8 wks; FU: 8 wks LF: 2 (2.6%) BC: no (1 sta- tistically sig- nificant differ- ence (% BSA higher in in- tervention group) Age: 48 Gender (per cent men)	Calcipotriol 50 mcg/g ointment BD, up to 120 g/ wk Placebo	Local AEs: not assessed Systemic AEs: blood and urine chem- istry analysis: parathy- roid hormone, serum calcium, bone-specific alkaline phos- phatase, urinary hyrox- yproline, 24- hr urinary cal- cium ex- cretion. Bone densitometry	No adverse ef- fects on bone metabolism or calcium	Bristol-Myers	Unclear

		to 20% EXCLU- SION CRITE- RIA: hypercal- caemia, bone, thyroid or parathyroid disease; topi- cal therapy within previ- ous 2 wks; systemic/ phototherapy within previ- ous 8 wks					
Heng 1990	DESIGN: be- tween-patient (retrospective study) patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: NA WITH- DRAWAL/ DROPOUT: NA	TD: 6 mths to 12 ys; FU: 6 mths to 12 years LF: NA BC: demo- graphic: yes; clinical: not demonstrated	nated steroids Con-	cal AEs: light/ electron mi- croscopy for examination of basal ker- atinocyte her- niation	vealed no be- tween-group	Sponsorship not reported Non- psoriatic con- trol group also considered 16/28 partici- pants had plaque psoria- sis	Not applicable

		(range: 6 mths to 12 years). Control group matched for age and gen- der EXCLU- SION CRI- TERIA: NR					
Katz 1987b	DESIGN: be- tween-patient patient deliv- ery ALLOCA- TION: random Method of randomisa- tion: NS Concealment: unclear BLIND- ING: double- blind (partici- pant/ investigator) WITH- DRAWAL/ DROPOUT: described	TD: 3 wks; FU: 3 wks LF: NA BC: demographic: yes; clinical: yes (gender imbalance) Age: 44 (range: 18 to 66) Gender (per cent men)	Betametha- sone dipropi- onate (0.05%) in opti- mised vehicle BD; Clobeta- sol 17 propi- onate (0.05%) ointment BD	assessed Systemic AEs: morning plasma corti- sol levels; 24-	adrenal axis in 8/40 partici-	not reported; 1 author from Schering Cor-	Unclear

		tion; hyper- sensitivity to study med- ications; con- current medi- cation that could af- fect study out- comes; use of systemic ther- apies within previ- ous 4 wks; use of topical ther- apies within previous 2 wks					
Katz 1989	DESIGN: within-patient patient deliv- ery ALLOCA- TION: random Method of randomisa- tion: unclear Concealment: unclear BLIND- ING: double- blind (partici- pant/ investigator) WITH- DRAWAL/ DROPOUT: described	LF: 0 (0%) BC: yes Age: 55 (range: 36 to 69) Gender (per cent men) : 53% Severity: NR INCLUSION CRITERIA:	Betametha- sone dipropi- onate (0.05%) in opti- mised vehicle BD (BMD) Clo- betasol propi- onate (0.05%) ointment BD (CP) Unin- volved (non- psoriatic) area used as test area for each participant	surface micro- scopic exami- na- tion with pho- tographic doc- umen-	served with ei- ther treat- ment. Preatro- phy identified in 20% of in- volved plaques (BMD: 11/ 59; CP: 12/ 59) and was more likely in females. In the test area (non- psoriatic skin) , 5% of plaques showed preat- rophy (BMD: 2/30; CP: 1/	Sponsored by Schering Cor- poration	Unclear
Kimball 2008 Phase II	DESIGN: un- controlled patient deliv- ery	N: 32 TD: 2 wks; FU: NS	Clobetasol 0. 05% foam BD Clo-	Maximal plasma concen-	Higher but non- significant lev-	Supported by Stiefel Labora- tories, Inc.	Not applicable

	ALLOCA- TION: unclear BLINDING: open WITH- DRAWAL/ DROPOUT: described	LF: 1 (3.1%) BC: NA Age: 24 to 72 Gender (per cent men): NS Severity: NS % white: 94% INCLUSION CRITERIA people with mild to mod- erate plaque psoriasis; aged > = 12 EXCLU- SION CRI- TERIA: NS	betasol 0.05% ointment BD (= 7 g/day, up to 50 g/wk)	tration of clo- betasol propi- onate		Review of phase II stud- ies on AD and psoriasis (clo- betasol foam)	
Kimball 2008 Phase III	DESIGN: be- tween-patient patient deliv- ery ALLOCA- TION: random Method of randomisa- tion: not stated Concealment: unclear BLIND- ING: double- blind (partici- pant/ investigator) WITH- DRAWAL/ DROPOUT: described	TD: 2 wks; FU: NS LF: 16 (3%) BC: NS Age: NS Gender (per cent men): NS Severity: most had baseline ISGA of 3 INCLUSION CRI- TERIA: peo- ple with mild	05% foam BD Clo- betasol 0.05% ointment BD Placebo foam BD Face, scalp, and inter- triginous areas	Treatment- related adverse events: • all • atrophy • burning • pruritis folliculitis ISGA (inves- tigator's static global assess- ment score) (scale NR)	(N = 253) C ointment: NS (N = 121) Placebo foam: 1% (N = 123) Burning: C foam: 2% (N = 253) C ointment:	Supported by Stiefel Labora- tories Inc. Re- view of phase III studies on AD and pso- riasis (clobeta- sol foam)	Unclear
Kragballe 1991b	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non-	BC: NA Age:	Cal- cipotriol oint- ment 50 mcg/ g BD (max: 100 g/wk)	Local AEs: pa- tient report of adverse events Investi- gator report of adverse events	AE(L): 3/15 (reported to be transient & mild).	Sponsorship not reported. Leo Pharma- ceuticals supplied study medication	Notapplicable

	random Method of randomisa- tion: NA Con- cealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	to 71) Gender (per cent men) : 53% Severity: % BSA: 14% (range: 5% to 30%) Most 'moder- ate' severity INCLUSION CRITE- RIA: partici- pants previ- ously respond- ing to cal- cipotriol dur- ing 8-wk clin- ical trial, but who had since relapsed EXCLU- SION CRITERIA: hyper- calcaemia, im- paired renal/ hepatic func- tion, daily re- ceiving > 400 i.u. vitamin D	No control	dermal atrophy. Systemic AEs: lab- oratory tests: FBC, serum	atrophy found in 4/8 partici- pants Systemic AEs: no consistent changes in lab- oratory analy- ses, with no clinically im- portant	treated with emollient or hydrocorti- sone cream 1% (not cal-	
Lambert 2002	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/	TD: 26 wks; FU: 26 wks	Tacalci- tol ointment, 4 mcg/g OD. No control	ticipants asked about adverse events. Tolera-	ment-related). AE(L): 26/ 157 (transient skin irritation) Tolerability at least moderate in 95% of par- ticipants Systemic AEs: no serious ad-	Hermal/BHI, Germany Scalp excluded Similar study to van de Kerkhof 2002, but un- clear whether Lambert 2002	Not applicable

	DROPOUT: described	plaque psoria- sis; aged 18 to 70; BSA 7% to 20%; labo- ratory param- eters normal at baseline EXCLU- SION CRI- TERIA: preg- nancy or risk thereof; topi- cal antipso- riatic therapy within previ- ous 2 wks; sys- temic antipso- riatic therapy within previ- ous 6 wks; retinoids within previ- ous 52 wks.; history of hy- perparathy- roidism; con- comitant use of drugs affecting cal- cium metabolism		bilirubin, creatinine, alkaline phosphatase, aspartate aminotrans- ferase, alanine aminotrans- ferase, gamma glutamyl- transpepti- dase, calcium, phosphate, sodium, potassium, glucose, urea, albumin. Urinalysis: calcium, creatinine, phosphate	calcaemia		
Lebwohl 1998b	DESIGN: be- tween-patient patient deliv- ery ALLOCA- TION: random. Method of randomisa- tion: unclear Concealment: unclear BLIND- ING: double- blind (partici- pant/ investigator)	TD: 26 wks;	imen: all par- ticipants received 2 wks of calcipotriol		Local AEs: AE(L) (treat- ment- related; all ir- ritant contact dermati- tis): Group A: 4/17 Group B: 1/20 No cuta- neous atrophy observed	Sponsored by Westwood Squibb Phar- maceuticals	Unclear

	WITH- DRAWAL/ DROPOUT: described	CRITERIA: people with at least moderate improve- ment in re- sponse to ini- tial 2-wk ther- apy regimen; aged ≥ 18 ; sta- ble dis- ease; BSA < = 20% (exclud- ing face/scalp) ; plaque eleva- tion at least moder- ate; willing to comply with study protocol EXCLU- SION CRITERIA: history of sen- sitivity to study ingre- dients; topical antipsoriat- ics within pre- vious 2 wks; UVB/PUVA within previ- ous 8 wks; his- tory of hyper- calcaemia, re- current illness	(weekends) Group B: Placebo oint- ment (weekdays) plus halo- betasol 0.05% ointment BD (weekends)				
Lebwohl 2001	DESIGN: be- tween-patient patient deliv- ery ALLOCA- TION: random Method of randomisa- tion: NR Concealment: unclear BLIND-	TD: 26 wks;	1% gel plus clo- betasol propi- onate 0.05% ointment for 6 wks. Once daily initially, then 'tapered'.	Local AEs: no. steroid- specific side- effects With- drawals due to adverse events (WA) Drug- related adverse events Systemic AEs: not assessed	effects WA: 0/50 AE(L) (treat- ment-related): TC: 24% TP:	Sponsorship: not reported No adequate effective- ness data re- ported. Num- bers of partic- ipants in each group NR	Unclear

	ING: double- blind (partici- pant/ investigator) WITH- DRAWAL/ DROPOUT: not described	participants participated in an open- label treat- ment phase for 6 wks (tazarotene gel 0.1% OM, clobetasol propionate ointment 0. 05% ON) EXCLU- SION CRI- TERIA: topi- cal antipsori- atic treatment within pre-	: tazarotene gel, 0.1%, OM (3/ 7 days) , plus clobeta- sol propionate 0.05% oint- ment ON (2/ 7days) (TC) Tazarotene gel, 0. 1%, OM (3/7 days), placebo ointment OM (2/7 days) , placebo oint- ment ON (2/7 days) (TP) Placebo gel OM (3/7 days), placebo ointment ON				
Menter 2007; Feldman 2007a	DESIGN: un- controlled patient deliv- ery ALLOCA- TION: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	TD: 4 wks; FU: 4 wks LF: 2 (0.1%) BC: NA Age: 49.7 (14. 7SD)	Clo- betasol 0.05% foam BD, up to 50 g/wk either as monother- apy or adjunc- tive to existing therapy	ing (0 none to	Erythema: 23. 7% peeling/scal- ing: 21.0% dryness: 28. 3% stinging/ burning: 15. 1% Telangiecta- sia, skin atro- phy, folliculi- tis: present in less than 1% of participants (findings not reported sepa-	(COBRA) Sponsorship	Not applicable

		erate to severe plaque psoria- sis; 3% to 20% BSA involve- ment EXCLU- SION CRI- TERIA: cur- rent systemic therapy, pho- tother- apy or topical therapy			rately) Pruritus: 5. 7%		
Miyachi 2002	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	TD: 54 wks; FU: 54 wks LF: 6 (3.8%) BC: NA Age: 48.2 (16. 1SD) Gender (per cent men) : 82% Severity:	Tacal- citol ointment 20 mcg/g OD (max: 10 g/ day) No control	treatment- related adverse events Sys- temic events: haematologi- cal tests (FBC) , blood bio- chemical tests (calcium, in- organic phos- phorus, albu- min, protein, bilirubin, urea nitrogen, crea- tinine, GP/ AST, GPT/ ALT, alkaline phosphatase, LDH, intact PTH), urinal-	mild to mod- erate) Systemic AEs: AE(S): 85/154 (155 events, of which 6 were considered treatment-re- lated). Serum levels of in- tact PTH and tacalcitol de- creased, sug- gesting percu- ta- neous absorp-	Scalp treated in 74/154 par- ticipants Usual dosing regimen for tacalcitol is	Not applicable

		riatics within previous 2 wks			High- dose tacalcitol affected serum cal- cium in par- ticipants with reduced renal function		
Poyner 1993	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	TD: 48 wks; FU: 48 wks LF: 59 (29. 1%) BC: NA Age: 43. 8 (range: 17 to 80)	Calcipotriol 50 mcg/g ointment No control	events (WA) Systemic AEs: biochemical and haemato- logical tests Compliance: self-reported	WA: 8/203 AE(L): 83/ 203 142 events re- ported by 83 (41%) partici- pants with 20. 2% being lesional/ perilesional ir- ritation Systemic AEs: no significant changes in	Leo Pharma- ceuticals Face/scalp ex-	Not applicable

		oral calcium/ vitamin D. topical an- tipsoriatics, lithium, sys- temic steroids				
Ramsay 1994	DESIGN: un- controlled open study patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	N: 167 TD: 52 wks; FU: 52 wks; FU: 52 wks LF: 39 (23.4%) BC: NA Age: 49 (range: 20 to 85) Gender (per cent men) : 60% Severity: PASI (modified): 8. 1 (6.7SD) INCLUSION CRITERIA: plaque psori- asis; previous response to calcipotriol; managed by specialists EX- CLUSION CRITERIA: pregnancy or risk thereof; abnormal serum calcium or phosphate; impaired hepatic/renal function; concomitant oral calcium/ vitamin D; systemic ther- apy within previous 8 wks; topi- cal therapy within previ- ous 4 wks	Calcipotriol 50 mcg/g ointment. Max dose: 100 g/wk; 2500 g/ pa Face/scalp/ neck excluded No control	counts) and biochemistry (bilirubin, AST/ALT, alkaline phos- phatase, albu- min, urate, creatinine, phosphate, to- tal calcium)	52/161 60 (46 considered to be treatment- related) events reported by 52 of 161 partic- ipants. 1 par- ticipant devel- oped a significant rise in serum cal- cium. No other ab- normal- ities in haema- tology or bio- chemistry tests. 118/161 participants reported con- tinuous medi- cation use and 80% to 90% used it twice daily. Mean use: 35. 1 g/wk ('ini- tially') to 23. 4 g/wk during	 Not applicable

Table 24.	Included	studies	of adverse events	(Continued)
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Roelofzen 2010	cer registry) and prospec- tive	TD: pix lithantracis (med): 4 mths (1 to 300 mths) liquor carbo- nis deter- gens (med): 6 mths (1 to 500 mths) FU: (med) 21 yrs LF: 329 (7.	apies. Focus of study is on coal tar: • Liquor • carbonis only (LCD) • Pix	malignancies Specific tu-	tion. How- ever, data on duration of tar	Not applicable-	Not applicable-
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van	DESIGN: un-	N: 58	Part 1: double-	Local AEs: oc-	Local AEs:	Sponsorship	Not applicable
de Kerkhof	controlled	TD: $< = 60$	blind study (8	currence of	WA: 0/58	not reported	
1997Ь	study	wks; FU: < =	wks): tacalci-	adverse events	AE		
	patient deliv-	60 wks	tol 4 mcg/g	(duration,	(L): 10/58 (19	Follow-up	
	ery	LF: 16 (27.	ointment OD	severity, and	events) AE(L)	study to Van	
	ALLOCA-	6%)	Placebo	whether treat-	(treatment-	de Kerkhof	
	TION: non-	BC: NA		ment-related)	related): 8/58	1996b - 3 of	
	random	Age:	Part 2: open	Participant	Toler-	15 centres par-	
	Method	45 (range: 19	follow-	and investiga-	ability: inves-	ticipated	
	of randomisa-	to 78)	up study (4 wk	tor assess-	tigator assess-		
	tion: NA Con-	Gender (per	wash-out pe-	ments of toler-	ment: 2.60 (0.	Scalp	
	cealment: NA	cent men): 69.	riod): tacalci-	ability (4-pt: v.	53SD, N = 58)	excluded	
	BLINDING:	0%	tol 4 mcg/g	good (3) to in-	;		
	open	Severity: BSA:	ointment OD,	sufficient (0))	participant as-		
	WITH-	8.6% (3.9SD)	< = 20 mg/day		sessment: 2.53		
	DRAWAL/	; TSS (0 to 12)	and < 2000	Systemic AEs:	(0.63SD, N =		
	DROPOUT:	:7.9 (2.1SD)	g per partici-	haematology	58)		
	described	INCLUSION	pant over	(erythrocytes,			
		CRITERIA:	study period.	platelets, hae-	Systemic AEs:		
		people with	Participants	moglobin,	AE(S): 0/58		
		chronic	could discon-	haematocrit)	No case of hy-		
		plaque pso-	tinue treat-	; blood chem-	percalcaemia		
		riasis par-	ment after 12	istry (serum			
		ticipating	wks	calcium, inor-			
		in previous		ganic phos-			
		double-blind	No control	phate, cre-			
		study (Van		atinine, ASAT,			
		de Kerkhof		alkaline phos-			
		1996b); aged		phatase,			
		25 to 80;		LDH)			
		normal serum					
		calcium/phos-					
		phate. EX-					
		CLUSION					
		CRITERIA:					
		pregnancy or					
		risk thereof;					
		topical ther-					
		apy within					
		previous 4					
		wks; systemic therapy					
		within previ-					
		ous 8 wks; se-					
		rious disease;					
		known allergy					
		to study					
		to study					

		medication; concomitant medication that could interfere with study drug or systemic calcium metabolism					
van de Kerkhof 2002c (see also Lambert 2002)	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	N: 304 TD: 13 wks; FU: 13 wks LF: 47 (15. 5%) BC: NA Age: 44 (range: 15 to 76) Gender (per cent men) : 57% Severity:	Tacal- citol 4 mcg/ g OD. Treat- ment discon- tinued during remission and restarted if re- lapse No control	Lo- cal AEs: num- ber treatment- related adverse events; with- drawals due to adverse events (WA); inves- tigator assess- ment of tol- erability; par- ticipant assess- ment of toler- ability Systemic AEs: Haema- tology: serum calcium, parathy- roid hormone (PTH), calci- tonin, cal- citriol Urine: calcium, crea- ti- nine, calcium/ creatinine ratio. Compli- ance with medication	AE(L): 65/ 304 Tol- erability excel- lent/good in 76% (patient assessment) to 92% (investi- gator assessment) of participants at final assessment. Systemic AEs: No clinically signif- icant changes in rou- tine haematol- ogy, urinalysis or serum chem- istry. Compli- ance with treatment reg- imen varied	Hermal/BHI, Germany Scalp	Not applicable

		cluding scalp); aged 18 to 70; normal base- line laboratory values Part 2 of study: respon- ders to part 1 (≥ 30% re- duction in sum score (TSS) from baseline) EXCLU- SION CRI- TERIA: top- ical steroids in previous 2 wks; systemic antipsoriat- ics within previous 6 wks; retinoids within previ- ous 52 wks; known hy- persensitivity to vitamin D serious con- comitant dis- ease; disease that might interfere with study assessments; concomitant use of oral calcium/ vitamin D; pregnancy or risk thereof			daily dose of 5 g (up to 13 g daily), but there was no effect on cal- cium home- ostasis. Dura- tion of excess dosing not re- ported		
Vazquez- Lopez 2004	DESIGN: un- controlled study patient deliv- ery	N: 20 TD: 26 wks; FU: 34 wks LF: 0 (0%) BC: NA	Clobeta- sol propionate 0.05% cream, OD (week-	Local AEs: clinical (naked eye) examina- tion of psori-	Overuse of topical steroids resulted in	Links compli- ance with ad- verse events	Not applicable

	ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	Age: 28. 2 (range: 20 to 55) Gender (per cent men) : 40% Severity: NR INCLUSION CRITERIA: absence of vis- i- ble or dermas- copic red lines (linear telang- iectasias) EXCLU- SION CRI- TERIA: use of topical steroids in pre- vious 2 mths	mcg/g ointment BD	and surround- ing area Dermo- scopic exami- nation of pso- riatic plaque	unapparent but dermo- scopically apparent linear telang- iectasias. 7/20 participants failed to adhere to recommended steroid dosing schedules. Dermoscopic	Study received no funding	
Veraldi 2006	DESIGN: un- controlled study patient deliv- ery ALLO- CATION: se- quential recruitment BLINDING: open WITH- DRAWAL/ DROPOUT:	TD: 45 dys; FU: 45 dys LF: 5 (10.4%) BC: NA	gel, short con- tact therapy (applied OD for 20 minutes	Burning (4-pt: 0 = absent to 3		Sponsorship not reported	Not applicable

	described	people with chronic stable plaque psoria- sis; % BSA < = 20% EXCLU- SION CRITERIA: use of antipso- riatic therapy within previ- ous 2 wks; concur- rent use of other top- ical, photo or systemic ther- apies			ticipant with- drew because of irritation on treated lesion		
Wishart 1994	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: groups deter- mined accord- ing to BSA af- fected. BLINDING: open WITH- DRAWAL/ DROPOUT: described	TD: 6 wks; FU: 6 wks LF: 1 (3%) BC: NA Age: 42.5 (13. 2SD) Gender (per cent men) : 47% Severity: Du- ration (mths): 202 (176SD) INCLUSION CRITERIA: people aged >18 with severe chronic plaque psoria- sis; le-	10): 8% to 15% BSA treated (600 to 1200 cm ²)	ECG Haematology, biochemistry, urine protein and glucose. Serum calcium, phos- phorus, plasma PTH, serum 25-hy- drox- yvitaim D, 1- alpha,25dihy- droxy-vi- taim D, 24 hr urine tests for calcium, creatinine and	306.1 mcg. No systemic adverse events, no skin irrita- tion. No clinically rel- evant changes in vital signs, haematol- ogy, biochem- istry, urine or ECGs IAGI (0 to 5): - 3.57 (1.01SD,	Usual dose is 3 mcg/ g BD, max. 30 g daily Sponsored by Solvay Duphar	Not applicable

psoriasis, con- current use of medicines contain- ing calcium or vitamin D, antacids or
digitalis

per cent men: per cent male; AE(L): number local adverse events/number participants; AE(S): number systemic adverse events/number participants; AE: adverse events; BC: baseline comparability; BD: twice daily; BSA: body surface area; FU: follow up (includes TD); N: number enrolled; NA: not applicable; NR: not reported; OD: once daily; PASI: Psoriasis Area and Severity Index; PRN: as required; TD: treatment duration; TSS: Total Severity Score; WA: withdrawal due to adverse events

Study	Reason for exclusion
Aste 2004	Follow-up under 12 wks and not focused on adverse events
Bos 2002	Not psoriasis, short review (letter)
Breneman 2007	Not a product included in our review (bexarotene gel 1%)
Carboni 2005	Not focused on adverse events
Feldman 2007a	Evaluated add-on clobetasol for participants treated concurrently with topical or systemic therapy
Floden 1975	Inadequate reporting of adverse events
Franssen 1999	Small (N = 54) retrospective study using participant questionnaires - aimed to identify teratogenetic effects of tar, but many women unable to recall whether tar used in pregnancy
Hong 2010; Hong 2011	Not chronic plaque psoriasis: paediatric dermatology participants had eczema or "eczema-psoriasis overlap (atopic dermatitis with associated features of psoriasis)"
Jacobi 2008	Small uncontrolled short-term and already reflected in results from main review
Kang 1998	Short-term and already reflected in results from main review
Lebwohl 1996	Follow-up under 12 wks and not focused on adverse events
Park 2002	Case study
Senter 1983	Adverse events not reported
Singh 2000	Short-term (4 weeks) and brief mention of adverse events

Table 25. Excluded studies of adverse events

Topical treatments for chronic plaque psoriasis (Review)

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Stevanovic 1977	Short-term, unclear if psoriasis, small numbers (N = 6)
Traulsen 2003	Participants were healthy volunteers
Uhoda 2003	Not about adverse events
Vissers 2004	Not about adverse events

Table 26. Included studies of compliance

Study	Methods	Participants	Interventions	Outcomes (compliance	Summary findings	Notes	Allocation concealment
Balkrishnan 2003	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: sin- gle-blind (par- ticipants un- aware of elec- tronic compli- ance assessment) WITH- DRAWAL/ DROPOUT: described	TD: 1 wk; FU: 1 wk LF: 0 (0%) BC: NA Age: NR Gender (per cent men) : NR Severity: NR INCLUSION CRI- TERIA: par- ticipants with psoriasis who already enrolling in a study with sal-	Topical sali- cylic acid 6% No control	Medication adherence: (1) MEMS cap: med- ication bottle cap with mi- croproces- sor to record time/date of every open- ing of the bot- tle. (2) Patient log (self report) of compliance Mean adher- ence rate: method 1: 67% (32% SD); method 2: 92% (7% SD)	(elec-	Sponsorship not reported	D
Carroll 2004a; Carroll 2004b; Carroll 2005	DESIGN: within-patient patient deliv- ery ALLOCA- TION:		Topical sal- icylic acid 6% plus 0. 1% tacrolimus ointment BD	adherence: (1) MEMS cap: med-	Adherence de- creased over time. On the interven- tion side, a de-	care, Inc. and	В

	random Method of randomisa- tion: NR Concealment: unclear BLINDING: Sin- gle-blind (par- ticipants un- aware of elec- tronic compli- ance assessment) WITH- DRAWAL/ DROPOUT: described	6 (range 18 to 70) Gender (per cent men) : 50% Severity: TSS (0 to 8): 5.3 INCLUSION CRI- TERIA: par- ticipants aged \geq 18; symmet- rical plaque- type psoriasis; BSA < = 10%; sym- metrical target plaque 1cm ² with each with a score of at least 1 for ery- thema, thick- ness, and scale EXCLU- SION CRI- TERIA: preg- nancy or risk thereof; topi-	Topical sal- icylic acid 6% plus placebo BD	time/date of every open- ing of the bot- tle.	herence rate of 10% was as- sociated with a 1-point in- crease in sever- ity (P < 0. 05). For the	sity School of Medicine. Excluded from effective- ness review	
		cal treatment within pre- vious 2 wks; phototherapy or sys- temic therapy within previ- ous 4 wks			% (doses taken/ doses ex- pected): 55%; % (days with twice- daily dose/to- tal days): 39.1% Higher adherence rate for women and older par- ticipants		
Feldman 2007	DESIGN: un- controlled study patient deliv- ery ALLOCA- TION: non-	TD: 8 wks; FU: 8 wks LF: NR BC: NA Age: 43.5	6% salicylic acid gel BD No control	its on partic- ipants' adher-	herence statis- tically signifi-	Sponsored in part by Astel- las Pharma US, Inc. The Center for Dermatol- ogy Research	D

	random Method of randomisa- tion: NA Concealment: NA BLINDING: sin- gle-blind (par- ticipants un- aware of elec- tronic compli- ance assessment) WITH- DRAWAL/ DROPOUT: described	(per cent men) : NR Severity: NR INCLUSION CRITERIA: NR EXCLU- SION CRI- TERIA: NR		Adherence as- sessed using MEMS cap: medica- tion bottle cap		is funded by a grant from GaldermaL- aboratories, LP. (see also Balkrishnan 2003; Carroll 2004a, 2004b, 2005)	
Ferrandiz 1998	DESIGN: be- tween-patient patient deliv- ery (therapy) Clinician delivery (pro- gramme) ALLOCA- TION: random Method of randomisa- tion: NR Concealment: unclear BLINDING: open WITH- DRAWAL/ DROPOUT: described	TD: 16 wks;	Calcipotriol plus reinforce- ment programme Cal- cipotriol with- out reinforce- ment programme	Reinforce- ment thera- peutic pro- gramme to en- hance adher- ence: derma- tologist provided par- ticipant edu- cation with explanation of disease charac- teristics and treatment effi- cacy and ap- plication, plus written infor- mation card	The reinforce- ment pro- gramme had no effect on treatment effi- cacy	Sponsorship not reported	В

		excipients; concurrent vi- ta- min D (> 400 units/day) or calcium tablets; psoria- sis mainly on face or hirsute areas					
Fouere 2005	DESIGN: questionnaire survey (obser- vational cross- sectional study) ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: response rate not reported	LF: NA BC: NA Age: 51.9 (SD 14.8)	Any antipsori- atic therapy	.1	rent treatment Main reasons for non-com- pliance: lack of ef- ficacy, messi- ness, and time constraints To im- prove compli- ance, patients suggested improved effi-	not reported. 70% of re- spon-	D
Gokdemir 2008	DESIGN: open uncon- trolled study patient deliv- ery ALLOCA- TION: non-	BC: NA Age:	scribed antip-	Medication ad- herence: num- ber prescribed doses taken/ number pre-	Mean adher- ence for topi- cal therapy: 72% (31%SD)	Findings relate to any treat- ment for pso- riasis (not just topical therapy)	D

	random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: described	to 70) Gender (per cent men) : 43% Sever- ity: PASI: 9.1 (range: 1.2 to 35) INCLUSION CRITE- RIA: chronic plaque psori- asis; received prescribed an- tipsoriatic therapy; aged \geq 16; attend- ing outpatient clinic in Istan- bul. EXCLU- SION CRI- TERIA: other types of psori- asis; hospi- talised; preg- nancy		scribed doses prescribed (see Zaghloul 2004).	Adherence rate was corre- lated with being un- married, more highly edu- cated, and be- ing satis- fied with treat- ment Main reasons for non-ad- herence were busyness and 'being fed up'	Sponsorship not reported	
Richards 1999	DE- SIGN: ques- tionnaire sur- vey (cross-sec- tional uncon- trolled study) patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: open WITH- DRAWAL/ DROPOUT: Response rate	Age: 49 (18 to 84) Gender (per cent men)	Any antipsori- atic therapy	Per cent com- plying with treatment (self report): scale not reported	39 per cent reported non- compliance (sometimes/ never comply- ing) with pre- scribed treat- ment. The non-compli- ant group had a higher self- rated disease severity, were younger, and had a younger age at onset. The non- compli- ant group re- ported that		D

Table 26.	Included	studies	of com	pliance	(Continued)
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	not reported	clinic; psoria- sis. EXCLU- SION CRITERIA: not stated			psoriasis had a greater impact on daily life Factors affect- ing compli- ance included the doctor- participant re- la- tionship; opti- mism with the treatment pre- scribed; and a lim- ited 'nuisance' value of treat- ment in terms of side-effects and hassle of use		
van de Kerkhof 1998	DE- SIGN: ques- tionnaire sur- vey (uncon- trolled study) patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: NA WITH- DRAWAL/ DROPOUT: Response rate reported	Severity: dura-	Any top- ical antipsori- atic therapy	Per cent com- ply- ing with fre- quency of ap- pli- cation of pre- scribed topical therapies Reason for non-com- pliance	of responders reported that the prescriber did not spec- ify dosage fre- quency. Where dosage frequency was specified, 33% (39%) complied with twice (once) daily regimens Main reasons for non-ad-	Psoriasis, the Journal of the Dutch Psoria- sis Patient Or- ganisation Responders asked to report on compli- ance over past	D

		TERIA: none stated					
van de Kerkhof 2000	DE- SIGN: ques- tionnaire sur- vey (uncon- trolled study) ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLINDING: single-blind WITH- DRAWAL/ DROPOUT: response rate reported	LF: NA Response rate: 14%	riatic therapy		ration of pre- scribed treat- ment (top- ical therapies): 71% Per cent com- ply- ing with fre- quency of ap- pli- cation of pre-	not reported 41- item question- naire mailed to 6100 sub- scribers of Psoriasis, the Journal of the Dutch Psoria- sis Patient Or- ganisation	D
van de Kerkhof 2001	DESIGN: within-patient (see Notes) patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA concealment: NA BLINDING:	FU: 8 wks LF: 93 (9.5%) BC: NR Age:	Calcipotriol cream OM plus cal- cipotriol oint- ment ON Calcipotriol ointment BD	Compliance: self-reported number of days cream/ ointment regi- men applied	men on most days. By wk 8, this statis- tic had fallen to 61% 51% of the 309 par-	not reported Control group comprised ret- rospective self- reported expe- rience of cal- cipotriol oint- ment monother-	D

	open WITH- DRAWAL/ DROPOUT: described	INCLUSION CRITE- RIA: psoriasis (type NR); eli- gible for treat- ment with cal- cipotriol EXCLU- SION CRITERIA: concomitant topical or sys- temic antipso- riatic therapy; co-exist- ing skin disor- der other than psoriasis				in the inter- vention group	
Zaghloul 2004	controlled study patient deliv- ery ALLOCA- TION: non- random Method of randomisa- tion: NA Concealment: NA BLIND- ING: single- blind (partici- pants unaware	N: 294 TD: 12 wks; FU: 12 wks LF: 93 (31. 6%) BC: NA Age: 45. 1 (range: 20 to 65) Gender (per cent men): 44. 3% Severity: NR INCLUSION CRITERIA: psoriasis (un- clear if chronic plaque only) ; aged 18 to 65; prescribed oral, topical or combined treatment EXCLU- SION CRI- TERIA: preg- nancy,	Topical, oral, or com- bined antipso- riatic medica- tion No control	30; higher score implies	Med- ication adher- ence measured by method 1 (ob- jective) much lower than by method 2 (pa- tient self report). Mean rate: 60.6% (33.0%SD) ; (range: 0% to169%) Direct correla- tion observed between med- ication adher- ence and qual- ity of life Adherence rate higher for par- ticipants who	Authors report no rel- evant financial interests	D

lactation, con- comitant dis- ease	were women, married, em- ployed, or not paying for pre- scriptions	
	Adherence greater for topical (vs. systemic) therapy, once daily, or first- time use	

per cent men: per cent male; AE(L): number local adverse events/number participants; AE(S): number systemic adverse events/number participants; AE: adverse events; BC: baseline comparability; BD: twice daily; BSA: body surface area; FU: follow up (includes TD); N: number enrolled; NA: not applicable; NR: not reported; OD: once daily; PASI: Psoriasis Area and Severity Index; PRN: as required; TD: treatment duration; TSS: Total Severity Score; WA: withdrawal due to adverse events

Table 27. Excluded studies of compliance

Study	Reason for exclusion
Atkinson 2004	Adherence not assessed
Chu 2000	Treatment guideline (not primary study)
Gupta 2007	Review/think piece
Lee 2006	Review
Osborne 2002	Study focused on non-responsive participants rather than those that are specifically non-compliant
Richards 2006	Review
Szeimies 2004	Think piece (not primary study)

APPENDICES

Appendix I. Specialised Register search strategy (RCTs)

Searched 04/12/04

Search updated on 17/11/2008: from January 2005 to date

Search updated on 3/02/2011: from November 2008 to date. One additional term added to 1st part: 'taclonex'.

Search updated 29/08/12: from February 2011 to date.

1st part

(*psorias* or *psoriat*) AND (*tar* or *gel* or alphosyl or carbodome or exorex or balneum or cocois or capasal or ceanel or ionil or meted or pentrax or anthralin or dithr* or micanol or psorin or psoriderm or salicylic or (vitamin and D) or calcipo* or dovo* or *calcit* or curatoderm or tazarotene or zorac or silkis or acitretin or neotigason or ciclosporin or cyclosporin or methotrexate or tacrolimus or pimecrolimus or protopic or elidel or retinoid* or macrolactam* or immunosuppressant* or taclonex) AND topical* 2nd part

(*psorias* or *psoriat*) AND ((adrenal and cortex and hormone*) or *steroid* or hydrocort* or cobadex or efcortelan or *derm or *dermal or *movate or mildison or calmurid or locoid or alclometasone or modrasone or beclo* or betametha* or betacap or betnovate or bettamousse or dipro* or clobetaso* or desox* or stiedex or diflucortolone or nerisone or fluocino* or synalar or metosyn or fluocortolone or ultralanum or flurandrenolone or fludroxycortide or haelan or fluticasone or cutivate or halci* or mometasone or elocon or triamcinolone or *cortyl)

Appendix 2. CENTRAL (Cochrane Library CD-ROM) and National Research Register (NRR) (CD-ROM interface) search strategies (RCTs)

CENTRAL (Cochrane Library CD-ROM 2005 issue 1): publication years 2001 to 2005-02-24 Searched on 17/11/08: CENTRAL (The Cochrane Library: http://www.thecochranelibrary.com/) Searched on 02/02/11: CENTRAL (The Cochrane Library: http://www.thecochranelibrary.com/) Searched on 23/08/12: CENTRAL (The Cochrane Library: http://www.thecochranelibrary.com/) National Research Register (CD-ROM interface, issue 2004/4): all projects with a start date of 2001 to 2005 Website browsed on 20/11/08: UK Clinical Research Network Study Portfolio (http://public.ukcrn.org.uk/search/) Website browsed on 10/09/12: UK Clinical Research Network Study Portfolio (http://public.ukcrn.org.uk/search/) Website browsed on 10/09/12: UK Clinical Research Network Study Portfolio (http://public.ukcrn.org.uk/search/)

- #1 MeSH descriptor Psoriasis explode all trees
- #2 (Psorias* or psoriat* or antipsoria*)
- #3 MeSH descriptor Coal Tar, this term only
- #4 "coal tar"
- #5 (alphosyl)
- #6 "carbo dome"
- #7 (clinitar or exorex or gelcosal or gelcotar)
- #8 (pragmatar or psorigel or balneum or polytar)
- #9 (psoriderm or tarcortin or cocois)
- #10 "T gel"
- #11 (capasal or ceanel or clinitar or ionil)
- #12 (meted or pentrax)
- #13 MeSH descriptor Anthralin, this term only
- #14 (Dithranol or dithrocream or micanol or psorin)
- #15 (salicylic next acid*)
- #16 (vitamin next d next analogue*)
- #17 (vitamin next d next derivative*)
- #18 (calcipotriol or calcipotriene or dovonex or dovobet)
- #19 (tacalcitol or curatoderm or tazarotene or zorac)
- #20 MeSH descriptor Calcitriol, this term only
- #21 (silkis or maxacalcitol)

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- #22 (#3 OR #4 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #
- 17 OR #18 OR #19 OR #20 OR #21)
- #23 MeSH descriptor Acitretin, this term only
- #24 MeSH descriptor Cyclosporins explode all trees
- #25 MeSH descriptor Methotrexate, this term only
- #26 MeSH descriptor Tacrolimus, this term only
- #27 (methotrexate or tacrolimus)
- #28 (pimecrolimus or elidel or protopic)
- #29 (acitretin or neotigason or cyclosporin or ciclosporin)
- #30 (#23 OR #24 OR #25 OR #26 OR #27 OR #28 OR #29)
- #31 (topical)
- #32 (#30 AND #31)
- #33 (topical next retinoid*)
- #34 (topical next macrolactam*)
- #35 (topical next immunosuppressant*)
- #36 MeSH descriptor Dermatologic Agents, this term only
- #37 MeSH descriptor Adrenal Cortex Hormones explode all trees
- #38 (corticosteroid* or (cortico next steroid*))
- #39 MeSH descriptor Hydrocortisone explode all trees
- #40 (hydrocortisone or cobadex or dioderm or efcortelan or hydrocortisyl mildison or alphaderm or calmurid)
- #41 "hydrocortisone butyrate"
- #42 "alclometasone dipropionate"
- #43 (modrasone or locoid or propaderm)
- #44 MeSH descriptor Beclomethasone, this term only
- #45 "beclomethasone dipropionate"
- #46 MeSH descriptor Betamethasone explode all trees
- #47 "betamethasone esters"
- #48 (betamethasone or betacap or betnovate or diprosone)
- #49 (diprosalic or bettamousse)
- #50 "clobetasol propionate"
- #51 "clobetasone butyrate"
- #52 (eumovate or trimovate or dermovate)
- #53 MeSH descriptor Desoximetasone, this term only
- #54 (desoxymethasone or desoximetasone or stiedex)
- #55 MeSH descriptor Diflucortolone, this term only
- #56 "diflucortolone valerate"
- #57 MeSH descriptor Fluocinolone Acetonide explode all trees
- #58 "fluocinolone acetonide"
- #59 (synalar or fluocinonide or metosyn or nerisone or elocon)
- #60 MeSH descriptor Fluocortolone explode all trees
- #61 (fluocortolone or ultralanum or flurandrenolone or haelan)
- #62 MeSH descriptor Flurandrenolone, this term only
- #63 (fludroxycortide)
- #64 (fluticasone next propionate)
- #65 (cutivate or halcinonide or halciderm)
- #66 "mometasone furoate"
- #67 MeSH descriptor Triamcinolone explode all trees
- #68 "triamcinolone acetonide"
- #69 (adcortyl or aureocort or nystadermal or triadcortyl)
- #70 MeSH descriptor Steroids explode all trees
- #71 (steroid*)
- #72 (#33 OR #34 OR #35 OR #36 OR #37 OR #38 OR #39 OR #40)
- #73 (#41 OR #42 OR #43 OR #44 OR #45 OR #46 OR #47 OR #48)

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- #74 (#49 OR #50 OR #51 OR #52 OR #53 OR #54 OR #55 OR #56)
- #75 (#57 OR #58 OR #59 OR #60 OR #61 OR #62 OR #63 OR #64)
- #76 (#65 OR #66 OR #67 OR #68 OR #69 OR #70 OR #71)
- #77 (#22 OR #32 OR #72 OR #73 OR #74 OR #75 OR #76)
- #78 ((#1 OR #2) AND #77), from 2008 to 2011

Appendix 3. MEDLINE (OVID) search strategy (RCTs)

MEDLINE (OvidSP Online http://www.ovid.com/): database updates 2002/07 to 2005/02 week 2

Search updated on 17/11/2008: 1950 to November Week 1 2008

Search updated on 22/02/2011: 1948 to January Week 3 2011

Search updated on 23/08/2012: 1946 to Present

- 1 randomized controlled trial.pt.
- 2 randomized controlled trial/
- 3 Random Allocation/
- 4 Double-Blind Method/
- 5 single-blind method/
- 6 clinical trial.pt.
- 7 exp clinical trial/
- 8 ((clinical\$ or intervention\$) adj5 (trial\$ or study or studies)).tw.
- 9 ((singl\$ or doubl\$ or trebl\$ or tripl\$) adj5 (blind\$ or mask\$)).tw.
- 10 Placebos/
- 11 (placebo\$ or random\$).tw.
- 12 Research Design/
- 13 Comparative Study/
- 14 exp evaluation studies as topic/
- 15 Follow-Up Studies/
- 16 Prospective Studies/
- 17 (control\$ or prospectiv\$ or volunteer\$).tw.
- 18 Animals/
- 19 Humans/
- 20 or/1-17
- 21 18 not (18 and 19)
- 22 20 not 21
- 23 exp Psoriasis/
- 24 psorias\$.tw.
- 25 psoriat\$.tw.
- 26 or/23-25
- 27 coal tar/
- 28 coal tar.tw.
- 29 alphosyl.tw.
- 30 carbo dome.tw.
- 31 clinitar.tw.
- 32 exorex.tw.
- 33 gelcosal.tw.
- 34 gelcotar.tw.
- 35 pragmatar.tw.
- 36 psorigel.tw.
- 37 balneum.tw.
- 38 polytar.tw.
- 39 psoriderm.tw.
- 40 tarcortin.tw.

41 cocois.tw. 42 (T adj gel).tw. 43 capasal.tw. 44 ceanel.tw. 45 clinitar.tw. 46 ionil.tw. 47 meted.tw. pentrax.tw. 48 49 Anthralin/ 50 dithranol.tw. 51 dithrocream.tw. 52 micanol.tw. psorin.tw. 53 54 salicylic acid\$.tw. 55 (vitamin adj d adj2 analogue\$).tw. 56 (vitamin adj d adj2 derivative\$).tw. 57 calcipotriol.tw. 58 calcipotriene.tw. 59 dovonex.tw. 60 dovobet.tw. tacalcitol.tw. 61 62 curatoderm.tw. 63 tazarotene.tw. 64 zorac.tw. 65 Calcitriol/ silkis.tw. 66 67 maxacalcitol.tw. 68 or/27-67 69 Acitretin/ 70 acitretin.tw. 71 neotigason.tw. 72 exp Cyclosporins/ 73 cyclosporin.tw. 74 ciclosporin.tw. 75 Methotrexate/ 76 methotrexate.tw. 77 Tacrolimus/ 78 tacrolimus.tw. 79 protopic.tw. 80 pimecrolimus.tw. 81 elidel.tw. 82 or/69-81 83 topical.tw. 84 82 and 83 85 topical retinoid\$.tw. 86 topical macrolactam\$.tw. 87 topical immunosuppressant\$.tw. 88 or/84-87 89 68 or 88 90 antipsoriat\$.tw. 91 antipsorias\$.tw. 92 26 or 90 or 91 93 exp Psoriasis/th, pc, dt [Therapy, Prevention & Control, Drug Therapy]

- 94 Dermatologic Agents/ 95 92 and 22 96 93 and 22 97 94 and 92 and 22 98 or/95-97 99 exp Adrenal Cortex Hormones/ 100 corticosteroid\$.tw. 101 cortico steroid\$.tw. 102 exp Hydrocortisone/ 103 hydrocortisone.tw.
- 104 cobadex.tw.
- 105 dioderm.tw.
- 106 efcortelan.tw.
- 107 hydrocortisyl.tw.
- 108 mildison.tw.
- 109 alphaderm.tw.
- 110 calmurid.tw.
- 111 hydrocortisone butyrate.tw.
- 112 locoid.tw.
- 113 alclometasone dipropionate.tw.
- 114 modrasone.tw.
- 115 Beclomethasone/
- 116 beclomet\$asone dipropionate.tw.
- 117 propaderm.tw.
- 118 exp Betamethasone/
- 119 betamethasone esters.tw.
- 120 betamethasone.tw.
- 121 betacap.tw.
- 122 betnovate.tw.
- 123 diprosone.tw.
- 124 diprosalic.tw.
- 125 bettamousse.tw.
- 126 clobetasol propionate.tw.
- 127 dermovate.tw.
- 128 clobetasone butyrate.tw.
- 129 eumovate.tw.
- 130 trimovate.tw.
- 131 Desoximetasone/
- 132 desoxymethasone.tw.
- 133 desoximetasone.tw.
- 134 stiedex.tw.
- 135 Diflucortolone/
- 136 diflucortolone valerate.tw.
- 137 nerisone.tw.
- 138 exp Fluocinolone Acetonide/
- 139 fluocinolone acetonide.tw.
- 140 synalar.tw.
- 141 fluocinonide.tw.
- 142 metosyn.tw.
- 143 exp Fluocortolone/
- 144 fluocortolone.tw.
- 145 ultralanum.tw.
- 146 Flurandrenolone/

- 147 flurandrenolone.tw.
- 148 fludroxycortid\$.tw.
- 149 haelan.tw.
- 150 fluticasone propionate.tw.
- 151 cutivate.tw.
- 152 halcinonide.tw.
- 153 halciderm.tw.
- 154 mometasone furoate.tw.
- 155 elocon.tw.
- 156 exp Triamcinolone/
- 157 triamcinolone acetonide.tw.
- 158 adcortyl.tw.
- aureocort.tw.
- 160 nystadermal.tw.
- 161 tri-adcortyl.tw.
- 162 exp Steroids/
- 163 steroid\$.tw.
- 164 or/99-163
- 165 22 and 92 and 164
- 166 22 and 92 and 89
- 167 98 or 165 or 166
- 168 (200811\$ or 200812\$).ed.
- 169 (2009\$ or 2010\$ or 2011\$).ed.
- 170 168 or 169
- 171 167 and 170

Appendix 4. EMBASE (OVID) search strategy (RCTs)

EMBASE (OvidSP Online http://www.ovid.com/): database updates 2002/08 to 2005/08

Search updated on 17/11/2008: 1980 to 2008 Week 46

Search updated on 31/01/2011: 1980 to 2011 Week 04

- Search updated on 23/08/2012: 1996 to 2012 Week 33
- 1 Coal Tar/
- 2 dithranol/ or tazarotene/ or 22 Oxacalcitriol/
- 3 (coal tar or alphosyl or carbo dome or clinitar or exorex or cocois or T gel or capasal or ceanel or ionil or meted or pentrax).ti,ab.
- 4 (gelcosal or gelcotar or pragmatar or psoriderm or psorigel or balneum or polytar or tarcortin or dithranol or dithrocream).ti,ab.
- 5 (anthralin or micanol or psorin or salicylic acid\$ or vitamin d analogue\$ or vitamin d derivative\$).ti,ab.
- 6 (calcipotriol or calcipotriene or dovonex or dovobet or tacalcitol or curatoderm or tazarotene or zorac or silkis or maxacalcitol).ti,ab.
- 7 vitamin d derivative/ or calcipotriol/ or calcitriol/ or tacalcitol/ or Salicylic Acid/
- 8 or/1-7
- 9 Etretin/ or Immunosuppressive Agent/ or Tacrolimus/
- 10 Cyclosporin/ or Retinoid/ or Pimecrolimus/
- 11 Methotrexate/

12 ((immunosuppressant\$ or acitretin or neotigason or cyclosporin\$ or ciclosporin\$ or methotrexate or retinoid\$ or macrolactam) adj5 topical\$).ti,ab.

- 13 ((tacrolimus or protopic or pimecrolimus or elidel) adj5 topical\$).ti,ab.
- 14 or/9-11
- 15 topical.ti,ab. or topical treatment/ or topical drug administration/
- 16 (14 and 15) or 12 or 13
- 17 exp Corticosteroid/
- 18 Hydrocortisone/
- 19 (corticosteroid\$ or cortico steroid\$ or hydrocortisone or cobadex or dioderm or efcortelan).ti,ab.

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- 20 (hydrocortisyl or mildison or alphaderm or calmurid).ti,ab.
- 21 (locoid or modrasone or beclomethasone dipropionate).ti,ab.
- 22 (alclometasone diproprionate or propaderm or betamethasone or betacap or betnovate or diprosone).ti,ab.
- 23 (diprosalic or bettamousse or clobetasol propionate or dermovate or clobetasone butyrate).ti,ab.
- 24 (eumovate or trimovate or desoxymethasone or desoxymethasone or desoximethasone or desoximethasone or stiedex).ti,ab.
- 25 (diflucortolone valerate or nerisone or fluocinolone acetonide or synalar or fluocinonide or metosyn).ti,ab.
- 26 (ultralanum or flurandrenolone or haelan or fluticasone propionate or cutivate or halcinonide or halciderm).ti,ab.
- 27 (mometasone furoate or elocon or adcortyl or aureocort or nystadermal or tri adcortyl or steroid\$).ti,ab.
- 28 beclometasone/ or psoralon/ or psoraderm/ or psoradexan/ or psorin/
- 29 Beclometasone Dipropionate/ or Urea/ or hydrocortisone butyrate/ or hydrocortisone plus urea/
- 30 alclometasone dipropionate/ or betamethasone dipropionate/ or betamethasone valerate/ or diflucortolone/

31 clobetasol propionate/ or clobetasone butyrate/ or desoximetasone/ or diflucortolone valerate/ or fluocinonide/ or Fluticasone Propionate/

- 32 fluocinolone/ or halcinonide/ or mometasone furoate.mp. or triamcinolone acetonide/
- 33 Fluocortolone/
- 34 Fludroxycortide/
- 35 Triamcinolone/
- 36 exp Steroid/
- 37 exp Steroid Hormone/
- 38 or/17-37
- 39 exp Antipsoriasis Agent/
- 40 8 or 16 or 38 or 39
- 41 exp Psoriasis/
- 42 (psorias\$ or psoriat\$ or antipsorias\$ or antipsoriat\$).ti,ab.
- 43 or/41-42
- 44 40 and 43
- 45 randomization/
- 46 Single Blind Procedure/
- 47 Double Blind Procedure/
- 48 exp clinical trial/
- 49 Placebo/
- 50 Methodology/
- 51 comparative study/
- 52 exp drug comparison/
- 53 evaluation/
- 54 Follow Up/
- 55 Prospective Study/
- 56 Crossover Procedure/
- 57 (clinical\$ adj3 (trial\$ or study or studies)).ti,ab.
- 58 (intervention\$ adj3 (trial\$ or study or studies)).ti,ab.
- 59 ((single\$ or doubl\$ or trebl\$ or tripl\$) adj3 blind\$).ti,ab.
- 60 ((single\$ or doubl\$ or trebl\$ or tripl\$) adj3 mask\$).ti,ab.
- 61 (placebo or placebos or random\$ or control\$ or prospectiv\$ or volunteer\$).ti,ab.
- 62 or/45-61
- 63 44 and 62
- 64 (2008\$ or 2009\$ or 2010\$ or 2011\$).em.
- 65 63 and 64
- 66 from 65 keep 1-2041

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Appendix 5. Science Citation Index (ISI web of Knowledge interface) and Conference Proceedings Citation Index - Science (ISI web of Knowledge interface) search strategies (RCTs)

Science Citation Index (SCI Web Of Knowledge http://wos.mimas.ac.uk/): publication years 2000 to 2005 Search updated on 17/11/2008: All lines linted as follows: DocType=All document types; Language=All languages; Database=SCI EXPANDED; Timespan=2005-2008

Search updated on 02/02/2011: All lines linted as follows: Databases=SCI-EXPANDED Timespan=2008-2011 Search updated on 23/08/12: (limited to 2011-01-01 - 2012-08-23)

Conference Proceedings Citation Index - Science (SCI Web Of Knowledge http://wos.mimas.ac.uk/)

Search updated on 02/02/2011: All lines linted as follows: Databases=CPCI-S Timespan=2008-2011

Search updated on 23/08/12: (limited to 2011-01-01 - 2012-08-23)

1 Topic=((psoria* OR antipsoria*) AND (trial* OR random* OR control OR controls OR double blind OR doubleblind or single blind OR singleblind OR placebo* or evaluation*))

2 Topic=(coaltar or coal tar or alphosyl or carbo dome or clinitar or exorex or gelcosal or gelcotar or pragmatar)

3 Topic=(psorigel or balneum or polytar or psoriderm or tarcortin or cocois or t gel or tgel or capasal or ceanel or clinitar or ionil or meted or pentrax)

4 Topic=(eumovate or trimovate or desoxime\$asone or desoxyme\$asone or stiedex or diflucortolone or nerisone or fluocinolone or synalar)

5 Topic=(anthralin OR dithranol or dithrocream or micanol or psorin or salicylic acid or salicylic acids or vitamin d or calcipotriol or calcipotriene or dovonex or dovobet or tacalcitol or curatoderm or tazarotene or zorac)

6 Topic=(fluocinonide or metosyn or fluocortolone or ultralanum or flurandrenolone or fludroxycortide or haelan or fluticasone propionate)

7 Topic=(calcitriol or silkis or maxacalcitol)

8 Topic=(cutivate or halcinonide or halciderm or mometasone furoate or elocon or triamcinolone or adcortyl or aureocort or nystadermal or triadcortyl or steroids or steroidal)

9 Topic=((acitretin or neotigason or cyclosporin* or ciclosporin* or methotrexate or tacrolimus or protopic or pimecrolimus or elidel or retinoid* or macrolactam* or immunosuppressant*) same topical)

10 Topic=(adrenal cortex hormone* or corticosteroid* or cortico steroid* or hydrocortisone or cobadex or dioderm or efcortelan or hydrocortisyl or mildison)

11 Topic=(alphaderm or calmurid or locoid or alclometasone dipropionate or modrasone or beclomet\$asone or propaderm)

12 Topic=(betamethasone or betacap or betnovate or diprosone or diprosalic or bettamousse or clobetasol propionate or dermovate or clobetasone butyrate)

13 #12 OR #11 OR #10 OR #9 OR #8 OR #7 OR #6 OR #5 OR #4 OR #3 OR #2

14 #13 AND #1

Appendix 6. BIOSIS (EDINAinterface) search strategy (RCTs)

BIOSIS (EDINAinterface): publication years 2001 to 2005

((((((((((((((()) (((() betamethasone or betacap or betnovate or diprosone or diprosalic or bettamousse or "clobetasol propionate" or dermovate or "clobetasone butyrate") and (sy: 2001-2005))) or ((al: alphaderm or calmurid or locoid or "alclometasone dipropionate" or modrasone or beclomethasone beclometasone or propaderm) and (sy: 2001-2005))) or ((al: "adrenal cortex hormone*" or corticosteroid* or "cortico steroid*" or hydrocortisone or cobadex or dioderm or efcortelan or hydrocortisyl or mildison) and (sy: 2001-2005))) or ((al: (acitretin or neotigason or cyclosporin* or ciclosporin* or methotrexate or tacrolimus or protopic or pimecrolimus or elidel or retinoid* or macrolactam* or immunosuppressant*)) and (sy: 2001-2005))) or ((al: cutivate or halcinonide or halciderm or "mometasone furoate" or elocon or triamcinolone or adcortyl or aureocort or nystadermal or triadcortyl or steroid or steroids or steroidal) and (sy: 2001-2005))) or ((al: (calcitriol or silkis or maxacalcitol)) and (sy: 2001-2005))) or ((al: fluocinonide or metosyn or fluocortolone or ultralanum or flurandrenolone or fludroxycortide or halean or "fluticasone propionate") and (sy: 2001-2005))) or ((al: anthralin OR dithranol or dithrocream or micanol or psorin or "salicylic acid" or "salicylic acids" or "vitamin d" or calcipotriol or calcipotriene or dovonex or dovobet or tacalcitol or curatoderm or tazarotene or zorac) and (sy: 2001-2005))) or ((al: (eumovate or trimovate or desoxime\$asone or desoxime\$asone or stiedex or diffucortolone or nerisone or fluocinolone or synalar)) and (sy: 2001-2005))) or (al: (calcitri or ionil or meted or pentrax))) or (al: (calcitra or or synalar)) and (sy: 2001-2005))) or (al: (calcitra or or sitelex or diffucortolone or nerisone or fluocinolone or synalar)) and (sy: 2001-2005))) or (al: (calcitra or desoxime\$asone or become or synalar)) and (sy: 2001-2005))) or (al: (calcitra or or sitelex or diffucortolone or nerisone or fluocinolone or synalar)) and (sy: 2001-2005))) or (al: (calcitra or cools or t gel o

blind" or doubleblind or "single blind" or singleblind or evaluation* or placebo* or control or controls) and (sy: 2001-2005)) and ((al: (psoria* or antipsoria*)) and (sy: 2001-2005)))))

Appendix 7. Dissertation Abstracts (Dialog Classic interface) and Inside Conferences (Dialog Classic interface) search strategies (RCTs)

Dissertation Abstracts (Dialog Classic interface): all publication years

Inside Conferences (Dialog Classic interface): all publication years

Biosis (Dialog Classic interface): all publication years

Search updated 17/11/2008: 1993-2008/Nov W2

Search updated 01/02/2011: 1993-2011/Jan W4

Search updated on 24/08/2012: 1993-2012/Aug W3

1 s (coal()tar or coaltar or alphosyl or carbo()dome or clinitar or exorex or cocois)/ti,ab,de

2 s (t()gel or tgel or capasal or ceanel or clinitar or ionil or meted or pentrax)/ti,ab,de

3 s (dithranol or gelcosal or gelcotar or pragmatar or psoriderm or psorigel or balneum or 4. polytar or tarcortin or dithrocream)/ ti,ab,de

4 s (micanol or psorin or salicylic()acid? ? or vitamin()d()analogue? or vitamin()d()derivative?)/ti,ab,de

5 s (calcipotriol or calcipotriene or dovonex or dovobet or tacalcitol or curatoderm or tazarotene or zorac)/ti,ab,de

6 s (calcitriol or silkis or maxacalcitol or adrenal()cortex()hormone? ?)/ti,ab,de

7 s ((cyclosporin or ciclosporin or methotrexate or acitretin or neotigason)(4w)topical?)/ti,ab,de

8 s ((retinoid? ? or immunosuppressant? ? or tacrolimus or protopic or pimecrolimus or macorlactam? ?)(4w)topical?)/ti,ab,de

9 s (corticosteroid? ? or cortico()steroid? ? or hydrocortisone or cobadex or dioderm or efcortelan)/ti,ab,de

10 s (hydrocortisyl or mildison or alphaderm or calmurid)/ti,ab,de

11 s (locoid or alclometasone()dipropionate or modrasone or beclomethasone()dipropionate)/ti,ab,de

12 s (propaderm or betamethasone or betacap or betnovate or diprosone)/ti,ab,de

13 s (diprosalic or bettamousse or clobetasol()propionate or dermovate or clobetasone()butyrate)/ti,ab,de

14 s (eumovate or trimovate or desoxymethasone or desoxymetasone or desoximetasone or desoximethasone or stiedex)/ti,ab,de

- 15 s (diflucortolone()valerate or nerisone or fluocinolone()acetonide or synalar or fluocinonide or metosyn)/ti,ab,de
- 16 s (fluocortolone or ultralanum or haelan or fluticasone()propionate or cutivate or halcinonide or halciderm or flurandrenolone or triamcinolone)/ti,ab,de

17 s (mometasone()furoate or elocon or adcortyl or aureocort or nystadermal or triadcortyl or steroid?? or steroidal)/ti,ab,de

- 18 s (beclometasone()dipropionate or fludroxycortide or triamcinolone)/ti,ab,de
- 19 s s1:s18
- 20 s (psoria? or antipsoria?)/ti,ab,de
- 21 s (random? or single()blind or singleblind or double()blind or doubleblind)/ti,ab,de
- 22 s (placebo or comparative()study or evaluation or prospective()study or crossover or trial or trials or triallist? ?)/ti,ab,de
- 23 s (control or controls? or prospectiv?)/ti,ab,de
- 24 s s21:s23
- 25 s s19 and s20 and s24
- 26 s UD=200812:201102
- 27 s s25 and s26
- 28 s rd s27

Appendix 8. SIGLE (WebSPIRS interface) search strategy (RCTs)

SIGLE (WebSPIRS interface): publication years 2001 to 2005 (database issue 2004/12)
SIGLE has not been updated since 2005, so searches were not rerun.
#1 psoriat* or psorias*(83 records)
#2 (2002 in PY) or (2003 in PY) or (2004 in PY) or (2005 in PY)(49881 records)
#3 ((2002 in PY) or (2003 in PY) or (2004 in PY) or (2005 in PY)) and (psoriat* or psorias*)(5 records)

Appendix 9. MEDLINE (OVID) search strategy (adverse events)

MEDLINE (OvidSP Online http://www.ovid.com/): database updates 1990 to 2005/02 week 2 Search updated on 17/11/08: 1950 to November Week 1 2008 Search updated on 02/02/11: 1948 to January Week 3 2011 Search updated on 23/08/12: 1946 to present

- 1 (coal adj tar).tw.
- 2 alphosyl.tw.
- 3 (carbo adj dome).tw.
- 4 clinitar.tw.
- 5 exorex.tw.
- 6 gelcosal.tw.
- 7 gelcotar.tw.
- 8 pragmatar.tw.
- 9 psorigel.tw.
- 10 balneum.tw.
- 11 polytar.tw.
- 12 psoriderm.tw.
- 13 tarcortin.tw.
- 14 cocois.tw.
- 15 (T adj gel).tw.
- 16 capasal.tw.
- 17 ceanel.tw.
- 18 ionil.tw.
- 19 meted.tw.
- 20 pentrax.tw.
- 21 dithranol.tw.
- 22 dithrocream.tw.
- 23 micanol.tw.
- 24 psorin.tw.
- 25 (salicylic adj acid\$).tw.
- 26 (vitamin adj d adj2 analogue\$).tw.
- 27 (vitamin adj d adj2 derivative\$).tw.
- 28 calcipotriol.tw.
- 29 calcipotriene.tw.
- 30 dovonex.tw.
- 31 dovobet.tw.
- 32 tacalcitol.tw.
- 33 curatoderm.tw.
- 34 tazarotene.tw.
- 35 zorac.tw.
- 36 silkis.tw.
- 37 maxacalcitol.tw.
- 38 antipsoriat\$.tw.
- 39 antipsorias\$.tw.

- 40 corticosteroid\$.tw.
- 41 (cortico adj steroid\$).tw.
- 42 hydrocortisone.tw.
- 43 cobadex.tw.
- 44 dioderm.tw.
- 45 efcortelan.tw.
- 46 hydrocortisyl.tw.
- 47 mildison.tw.
- 48 alphaderm.tw.
- 49 calmurid.tw.
- 50 (hydrocortisone adj butyrate).tw.
- 51 locoid.tw.
- 52 (alclometasone adj dipropionate).tw.
- 53 modrasone.tw.
- 54 (beclomet\$asone adj dipropionate).tw.
- 55 propaderm.tw.
- 56 (betamethasone adj esters).tw.
- 57 betamethasone.tw.
- 58 betacap.tw.
- 59 betnovate.tw.
- 60 diprosone.tw.
- 61 diprosalic.tw.
- 62 bettamousse.tw.
- 63 (clobetasol adj propionate).tw.
- 64 dermovate.tw.
- 65 (clobetasone adj butyrate).tw.
- 66 eumovate.tw.
- 67 trimovate.tw.
- 68 desoxymethasone.tw.
- 69 desoximetasone.tw.
- 70 stiedex.tw.
- 71 (diflucortolone adj valerate).tw.
- 72 nerisone.tw.
- 73 (fluocinolone adj acetonide).tw.
- 74 synalar.tw.
- 75 fluocinonide.tw.
- 76 metosyn.tw.
- 77 fluocortolone.tw.
- 78 ultralanum.tw.
- 79 flurandrenolone.tw.
- 80 haelan.tw.
- 81 (fluticasone adj propionate).tw.
- 82 cutivate.tw.
- 83 halcinonide.tw.
- 84 halciderm.tw.
- 85 (mometasone adj furoate).tw.
- 86 elocon.tw.
- 87 (triamcinolone adj acetonide).tw.
- 88 adcortyl.tw.
- 89 aureocort.tw.
- 90 nystadermal.tw.
- 91 tri-adcortyl.tw.
- 92 steroid\$.tw.

- 93 Coal Tar/
- 94 Anthralin/
- 95 Calcitriol/
- 96 exp Cyclosporins/
- 97 Tacrolimus/
- 98 Dermatologic Agents/
- 99 exp Adrenal Cortex Hormones/
- 100 exp Hydrocortisone/
- 101 Beclomethasone/
- 102 exp Betamethasone/
- 103 Desoximetasone/
- 104 Diflucortolone/
- 105 exp Fluocinolone Acetonide/
- 106 Fluocortolone/
- 107 Flurandrenolone/
- 108 Flurandrenolone/
- 109 exp Triamcinolone/
- 110 exp Steroids/
- 111 or/1-110
- 112 Acitretin/
- 113 Immunosuppressive Agents/
- 114 Cyclosporine/
- 115 Retinoids/
- 116 Methotrexate/
- 117 or/112-116
- 118 topical.ti,ab. or topical treatment/ or topical drug administration/
- 119 117 and 118

120 ((tacrolimus or protopic or pimecrolimus or elidel or immunosuppressant\$ or acitretin or neotigason or cyclosporin\$ or ciclosporin\$ or methotrexate or retinoid\$ or macrolactam) adj5 topical\$).tw.

- 121 119 or 120 or 111
- 122 (safe or safety).tw.
- 123 side effect\$.tw.
- 124 treatment emergent.tw.
- 125 undesirable effect\$.tw.
- 126 tolerability.tw.
- 127 toxicity.tw.
- 128 adrs.tw.
- 129 (adverse adj3 (effect or effects or reaction or reactions or event or events or outcome or outcomes)).tw.
- 130 Adverse Drug Reaction Reporting Systems/
- 131 drug hypersensitivity/
- 132 hypersensit\$.tw.
- 133 harm\$.tw.
- 134 exp Substance Withdrawal Syndrome/ci [Chemically Induced]
- 135 rebound.tw.
- 136 Hypercalcemia/ci [Chemically Induced]
- 137 exp Urinary Calculi/ci [Chemically Induced]
- 138 Tachyphylaxis/ci, de [Chemically Induced, Drug Effects]
- 139 exp Substance Withdrawal Syndrome/ci [Chemically Induced]
- 140 exp Atrophy/ci [Chemically Induced]
- 141 exp Telangiectasis/ci [Chemically Induced]
- 142 cutaneous atrophy.tw.
- 143 striae.tw.
- 144 skin atrophy.tw.

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- 145 exp Abnormalities, Drug-Induced/
- 146 exp Drug Toxicity/
- 147 or/122-146
- 148 Coal Tar/ae [Adverse Effects]
- 149 Anthralin/ae [Adverse Effects]
- 150 Calcitriol/ae [Adverse Effects]
- 151 Acitretin/ae [Adverse Effects]
- 152 exp Cyclosporins/de, ae [Drug Effects, Adverse Effects]
- 153 Methotrexate/ae [Adverse Effects]
- 154 Tacrolimus/ae [Adverse Effects]
- 155 Dermatologic Agents/ae [Adverse Effects]
- 156 exp Adrenal Cortex Hormones/ae, de [Adverse Effects, Drug Effects]
- 157 exp Hydrocortisone/ae [Adverse Effects]
- 158 Beclomethasone/ae [Adverse Effects]
- 159 exp Betamethasone/ae [Adverse Effects]
- 160 Desoximetasone/ae [Adverse Effects]
- 161 Diflucortolone/ae [Adverse Effects]
- 162 exp Fluocinolone Acetonide/ae [Adverse Effects]
- 163 Fluocortolone/ae [Adverse Effects]
- 164 Flurandrenolone/ae [Adverse Effects]
- 165 exp Triamcinolone/ae [Adverse Effects]
- 166 exp Steroids/ae [Adverse Effects]
- 167 or/148-166
- 168 147 or 167
- 169 (psorias\$ or psoriat\$).tw.
- 170 exp psoriasis/
- 171 or/169-170
- 172 121 and 168 and 171
- 173 exp animals/ not (exp animals/ and humans/)
- 174 172 not 173
- 175 (comment or editorial).pt.
- 176 174 not 175
- 177 (2008\$ or 2009\$ or 2010\$ or 2011\$).ed.
- 178 176 and 177

Appendix 10. EMBASE (OVID) search strategy (adverse events)

EMBASE (OvidSP Online http://www.ovid.com/): 1990 to 2005/08

Search updated on 11/12/08: 1980 to 2008 Week 49

Search updated on 02/02/11: 1980 to 2011 week 4

Search updated on 23/08/12: 1996 to 2012 Week 34

Note: A pragmatic approach was taken to reduce irrelevant records and to negate the over indexing of records in EMBASE; EMTREE terms were focused in this strategy

- 1 (coal adj tar).ti,ab.
- 2 alphosyl.ti,ab.
- 3 (carbo adj dome).ti,ab.
- 4 clinitar.ti,ab.
- 5 exorex.ti,ab.
- 6 gelcosal.ti,ab.
- 7 gelcotar.ti,ab.
- 8 pragmatar.ti,ab.
- 9 psorigel.ti,ab.

- 10 balneum.ti,ab.
- 11 polytar.ti,ab.
- 12 psoriderm.ti,ab.
- 13 tarcortin.ti,ab.
- 14 cocois.ti,ab.
- 15 (T adj gel).ti,ab.
- 16 capasal.ti,ab.
- 17 ceanel.ti,ab.
- 18 ionil.ti,ab.
- 19 meted.ti,ab.
- 20 pentrax.ti,ab.
- 21 dithranol.ti,ab.
- 22 dithrocream.ti,ab.
- 23 micanol.ti,ab.
- 24 psorin.ti,ab.
- 25 (salicylic adj acid\$).ti,ab.
- 26 (vitamin adj d adj2 analogue\$).ti,ab.
- 27 (vitamin adj d adj2 derivative\$).ti,ab.
- 28 calcipotriol.ti,ab.
- 29 calcipotriene.ti,ab.
- 30 dovonex.ti,ab.
- 31 dovobet.ti,ab.
- 32 tacalcitol.ti,ab.
- 33 curatoderm.ti,ab.
- 34 tazarotene.ti,ab.
- 35 zorac.ti,ab.
- 36 silkis.ti,ab.
- 37 maxacalcitol.ti,ab.
- 38 antipsoriat\$.ti,ab.
- 39 antipsorias\$.ti,ab.
- 40 corticosteroid\$.ti,ab.
- 41 (cortico adj steroid\$).ti,ab.
- 42 hydrocortisone.ti,ab.
- 43 cobadex.ti,ab.
- 44 dioderm.ti,ab.
- 45 efcortelan.ti,ab.
- 46 hydrocortisyl.ti,ab.
- 47 mildison.ti,ab.
- 48 alphaderm.ti,ab.
- 49 calmurid.ti,ab.
- 50 (hydrocortisone adj butyrate).ti,ab.
- 51 locoid.ti,ab.
- 52 (alclometasone adj dipropionate).ti,ab.
- 53 modrasone.ti,ab.
- 54 (beclomet\$asone adj dipropionate).ti,ab.
- 55 propaderm.ti,ab.
- 56 (betamethasone adj esters).ti,ab.
- 57 betamethasone.ti,ab.
- 58 betacap.ti,ab.
- 59 betnovate.ti,ab.
- 60 diprosone.ti,ab.
- 61 diprosalic.ti,ab.
- 62 bettamousse.ti,ab.

- 63 (clobetasol adj propionate).ti,ab.
- 64 dermovate.ti,ab.
- 65 (clobetasone adj butyrate).ti,ab.
- 66 eumovate.ti,ab.
- 67 trimovate.ti,ab.
- 68 desoxymethasone.ti,ab.
- 69 desoximetasone.ti,ab.
- 70 stiedex.ti,ab.
- 71 (diflucortolone adj valerate).ti,ab.
- 72 nerisone.ti,ab.
- 73 (fluocinolone adj acetonide).ti,ab.
- 74 synalar.ti,ab.
- 75 fluocinonide.ti,ab.
- 76 metosyn.ti,ab.
- 77 fluocortolone.ti,ab.
- 78 ultralanum.ti,ab.
- 79 flurandrenolone.ti,ab.
- 80 haelan.ti,ab.
- 81 (fluticasone adj propionate).ti,ab.
- 82 cutivate.ti,ab.
- 83 halcinonide.ti,ab.
- 84 halciderm.ti,ab.
- 85 (mometasone adj furoate).ti,ab.
- 86 elocon.ti,ab.
- 87 (triamcinolone adj acetonide).ti,ab.
- 88 adcortyl.ti,ab.
- 89 aureocort.ti,ab.
- 90 nystadermal.ti,ab.
- 91 tri-adcortyl.ti,ab.
- 92 steroid\$.ti,ab.
- 93 *Coal Tar/
- 94 *alphosyl/
- 95 *carbo dome/
- 96 *Salicylic Acid/
- 97 *capasal/
- 98 *meted/
- 99 *Dithranol/
- 100 *Psorin/
- 101 *Vitamin d Derivative/
- 102 *Calcipotriol/
- 103 *Betamethasone Dipropionate Plus Calcipotriol/
- 104 *Tacalcitol/
- 105 *Tazarotene/
- 106 *Calcitriol/
- 107 *22 Oxacalcitriol/
- 108 *Corticosteroid/
- 109 *Hydrocortisone/
- 110 *Urea/
- 111 *Hydrocortisone Butyrate/
- 112 *Alclometasone Dipropionate/
- 113 *Beclometasone Dipropionate/
- 114 *Betamethasone/
- 115 *Betamethasone Valerate/

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- 116 *Betamethasone Dipropionate/
- 117 *Clobetasol Propionate/
- 118 *Clobetasone Butyrate/
- 119 *trimovate/
- 120 *Desoximetasone/
- 121 *Diflucortolone Valerate/
- 122 *Fluocinolone Acetonide/
- 123 *Fluocinonide/
- 124 *Fluocortolone/
- 125 *Fludroxycortide/
- 126 *Fluticasone Propionate/
- 127 *Halcinonide/
- 128 *Mometasone Furoate/
- 129 *Triamcinolone Acetonide/
- 130 *Triamcinolone/
- 131 *Mycolog/
- 132 *exp Steroid/
- 133 *Cyclosporin Derivative/ or *Cyclosporin/
- 134 *Tacrolimus/
- 135 *Dermatological Agent/
- 136 or/1-135
- 137 *Tacrolimus/
- 138 *Pimecrolimus/
- 139 *Immunosuppressive Agent/
- 140 *Etretin/
- 141 *Cyclosporin/
- 142 *Cyclosporin A/
- 143 *Methotrexate/
- 144 *Retinoid/
- 145 or/137-144
- 146 topical.ti,ab. or topical treatment/ or topical drug administration/
- 147 145 and 146
- 148 ((tacrolimus or protopic or pimecrolimus or elidel or immunosuppressant\$ or acitretin or neotigason or cyclosporin\$ or ciclosporin\$ or methotrexate or retinoid\$ or macrolactam) adj5 topical\$).ti,ab.
- 149 136 or 147 or 148
- 150 (safe or safety).ti,ab.
- 151 side effect\$.ti,ab.
- 152 treatment emergent.ti,ab.
- 153 undesirable effect\$.ti,ab.
- 154 tolerability.ti,ab.
- 155 toxicity.ti,ab.
- 156 adrs.ti,ab.
- 157 (adverse adj3 (effect or effects or reaction or reactions or event or events or outcome or outcomes)).ti,ab.
- 158 *Safety/ or *Drug Safety/
- 159 *Side Effect/
- 160 *Adverse Drug Reaction/
- 161 *Drug Tolerability/
- 162 *Toxicity/ or *Drug Toxicity/
- 163 *Drug Surveillance Program/
- 164 *Adverse Outcome/
- 165 hypersensit\$.ti,ab.
- 166 harm\$.ti,ab.
- 167 rebound.ti,ab.

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- 168 *Drug Hypersensitivity/
- 169 *Rebound/
- 170 *Withdrawal Syndrome/
- 171 *Hypercalcemia/
- 172 *Urolithiasis/
- 173 *Tachyphylaxis/
- 174 *Drug Withdrawal/
- 175 *Atrophy/
- 176 *Telangiectasia/
- 177 cutaneous atrophy.ti,ab.
- 178 striae.ti,ab.
- 179 skin atrophy.ti,ab.
- 180 *Skin Atrophy/
- 181 *Stria/
- 182 or/150-181
- 183 *Coal Tar/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 184 *alphosyl/ae [Adverse Drug Reaction]
- 185 *Salicylic Acid/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 186 *Dithranol/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 187 *Psorin/ae [Adverse Drug Reaction]
- 188 *Vitamin d Derivative/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 189 *Calcipotriol/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 190 *Betamethasone Dipropionate Plus Calcipotriol/to, ae [Drug Toxicity, Adverse Drug Reaction]
- 191 *Tacalcitol/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 192 *Tazarotene/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 193 *Calcitriol/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 194 *22 Oxacalcitriol/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 195 *Corticosteroid/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 196 *Hydrocortisone/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 197 *Urea/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 198 *Hydrocortisone Butyrate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 199 *Alclometasone Dipropionate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 200 *Beclometasone Dipropionate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 201 *Betamethasone/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 202 *Betamethasone Valerate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 203 *Betamethasone Dipropionate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 204 *Clobetasol Propionate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 205 *Clobetasone Butyrate/ae [Adverse Drug Reaction]
- 206 *trimovate/ae [Adverse Drug Reaction]
- 207 *Desoximetasone/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 208 *Diflucortolone Valerate/to, ae [Drug Toxicity, Adverse Drug Reaction]
- 209 *Fluocinolone Acetonide/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 210 *Fluocinonide/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 211 *Fluocortolone/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 212 *Fludroxycortide/ae [Adverse Drug Reaction]
- 213 *Fluticasone Propionate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 214 *Halcinonide/ae [Adverse Drug Reaction]
- 215 *Mometasone Furoate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 216 *Triamcinolone Acetonide/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 217 *Triamcinolone/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 218 *Mycolog/ae [Adverse Drug Reaction]
- 219 *exp Steroid/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 220 *Cyclosporin Derivative/ae, to [Adverse Drug Reaction, Drug Toxicity]

Topical treatments for chronic plaque psoriasis (Review)

- 221 *Cyclosporin/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 222 *Tacrolimus/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 223 *Dermatological Agent/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 224 *Pimecrolimus/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 225 *Immunosuppressive Agent/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 226 *Etretin/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 227 *Cyclosporin A/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 228 *Methotrexate/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 229 *Retinoid/ae, to [Adverse Drug Reaction, Drug Toxicity]
- 230 or/183-229
- 231 182 or 230
- 232 (psorias\$ or psoriat\$).ti,ab.
- 233 *exp Psoriasis/
- 234 232 or 233
- 235 149 and 231 and 234
- 236 animal/ not (animal/ and human/)
- 237 235 not 236
- 238 ("200800" or "200900").em.
- 239 (2010\$ or 2011\$).em.
- 240 238 or 239
- 241 237 and 240

Appendix 11. EMBASE (OVID) search strategy (compliance)

EMBASE (OvidSP Online http://www.ovid.com/): 1980 to 2007 Week 49 Not updated in 2011

- 1 compliance\$.ti,ab. (46694)
- 2 complied.ti,ab. (1455)
- 3 compliance/ (1674)
- 4 comply.ti,ab. (3613)
- 5 (medicat\$ adj4 adher\$).ti,ab. (1732)
- 6 (drug\$ adj4 adher\$).ti,ab. (777)
- 7 (medicine\$ adj4 adher\$).ti,ab. (62)
- 8 (treatment adj4 adher\$).ti,ab. (3072)
- 9 concordance\$.ti,ab. (12825)
- 10 or/1-9 (68155)
- 11 psor\$.ti,ab. (20549)
- 12 10 and 11 (171)
- 13 from 12 keep 1-171 (171)

Appendix 12. MEDLINE (OVID) search strategy (compliance)

MEDLINE(OvidSP Online http://www.ovid.com/): 1950 to November Week 2 2007 Not updated in 2011 1. compliance\$.ti,ab. (52891) 2. complied.ti,ab. (1748) 3. compliance/ (2982) 4. comply.ti,ab. (4307)

- 5. (medicat\$ adj4 adher\$).ti,ab. (2047)
- 6. (drug\$ adj4 adher\$).ti,ab. (878)
- 7. (medicine\$ adj4 adher\$).ti,ab. (76)
- 8. (treatment adj4 adher\$).ti,ab. (3442)

9. concordance\$.ti,ab. (14973) 10. or/1-9 (78297) 11. psor\$.ti,ab. (25210) 12. 10 and 11 (162)

WHAT'S NEW

Last assessed as up-to-date: 2 February 2011.

Date	Event	Description
9 December 2015	Amended	Author information (affiliation) updated

HISTORY

Protocol first published: Issue 4, 2004 Review first published: Issue 2, 2009

Date	Event	Description
2 April 2013	Amended	'Declarations of interest' section updated.
25 February 2013	New citation required but conclusions have not changed	This review was updated, but there was no change to the conclusions
25 February 2013	New search has been performed	48 new RCTs added. Comparisons revised to reflect new evidence. Scalp trials moved to separate comparison groups ready for removal at next update (#18; #19). Inverse psoriasis trials also moved to separate compar- ison groups (#16; #17). Nail trials removed, as now part of separate Cochrane review
15 December 2008	Amended	Review amended to reflect peer review comments
25 June 2008	Amended	Converted to new review format.
29 September 2003	New citation required and conclusions have changed	Substantive amendment

Topical treatments for chronic plaque psoriasis (Review)

CONTRIBUTIONS OF AUTHORS

The following contributions were made by the authors stated. Link with editorial base and co-ordinate contributions from co-authors (AM). Draft protocol (AM with contributions from MC, GD, GE, and JM). Run searches (adherence) (AM). Identify relevant titles and abstracts from searches, i.e. broad screen (AM and JM). Obtain copies of trials (AM). Select which trials to include (AM, JM, and MC as arbitrator when necessary). Extract data from trials (AM, JM, and HH). Enter data into RevMan (AM). Carry out analysis (AM and JM) Interpret analysis (AM, JM, and HH). Draft final review (AM with contribution from MC, GD, HH, and JM). Update review (AM, JM, HH, and MC).

DECLARATIONS OF INTEREST

Mike Cork: "I gave a lecture for Leo Pharmaceuticals about psoriasis and atopic eczema in 2012."

Anne R Mason: none declared

Gordon Dooley: none declared

James Mason: none declared

Helen Hancock: none declared

The clinical referee for this review, Dr Phyllis Spuls, stated the following potential conflict of interest on her comments form: "I have participated in an advisory board of LEO Pharma to give guidance to general practitioners regarding what to do if patients used calcipotriol, which has been removed from the market. I am involved as a Principal Investigator in many clinical trials with systemic agents for psoriasis and now in one for the improvement of adherence to Dovobet."

SOURCES OF SUPPORT

Internal sources

- Funding from Centre for Reviews and Dissemination to update review (2002) for UK products only, UK.
- Award from University of York Fund for Staff on Fixed-Term Contracts, UK.

External sources

- Grant from Crookes Healthcare Ltd. to do original systematic review (1999), UK.
- Grant from the Psoriasis Association to update review (2011), Other.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

There are some differences between the protocol and the review.

1. In our protocol, we stated our intention to adjust for the precision of findings from within-patient studies for within-patient correlation. However, we were unsuccessful in our attempts to identify estimates of this correlation from published or unpublished sources. We therefore undertook sensitivity analysis to investigate differences between within-patient and between-patient studies.

2. In our protocol, we listed three primary outcome measures for data extraction. In the review, we also included the Patient Assessment of Global Improvement.

3. In our protocol, we stated that there would be no language restrictions when searching for publications. However, the search for longer-term studies of adverse events included a restriction to publications in English.

4. In our protocol, we stated our intention that studies meeting only some of the inclusion criteria stated above would be listed as excluded studies. However, as large numbers of studies would need to be listed, this was not feasible. Therefore, we listed as excluded studies only those studies that we deemed potentially eligible for inclusion *and* for which we retrieved full papers, but which we subsequently found to fail to meet the inclusion criteria.

5. Throughout the text, we replaced all references to vitamin D3 with 'vitamin D analogues'.

6. Under Types of studies, we relaxed the condition that studies were of at least two weeks duration. In the same section, we added the following sentence: "If no useful effectiveness, withdrawal or adverse events data were available, either from the published paper or from sponsors or trialists, we excluded the study."

7. Under Types of interventions, we added the sentence "The potency of topical corticosteroids was based on classifications from a previous review (Mason 2002b)".

8. Under Methods, Selection of studies, and our explanations of the studies excluded, we added the searches for studies exploring adverse events and compliance studies.

9. Under Methods/Data extraction and management, we added the phrase 'between-patient design'.

10. Under Methods/Assessment of risk of bias in included studies, we removed the phrase 'in each arm'.

11. Under Methods/Unit of analysis issues/'Summarising primary outcomes with standardised mean differences', we revised the text in this section to include the PAGI outcome and to provide a fuller explanation of our analytic approach.

12. Under Methods/Unit of analysis issues/Secondary outcomes, we gave an explanation regarding the different method of analysis used.

13 Under Methods/Data collection and analysis/Sensitivity analysis, we added text to describe the different types of sensitivity analysis undertaken.

14 We replaced 'standardised weighted mean difference' with 'standardised mean difference' throughout the text to reflect Cochrane terminology.

15. Under Methods/Data and analyses/Subgroup analysis and investigation of heterogeneity, we inserted two paragraphs to explain our approach to statistical heterogeneity.

Topical treatments for chronic plaque psoriasis (Review)

INDEX TERMS

Medical Subject Headings (MeSH)

Administration, Topical; Adrenal Cortex Hormones [adverse effects; *therapeutic use]; Bone Density Conservation Agents [adverse effects; *therapeutic use]; Chronic Disease; Facial Dermatoses [drug therapy]; Psoriasis [*drug therapy]; Randomized Controlled Trials as Topic; Scalp Dermatoses [drug therapy]; Vitamin D [adverse effects; analogs & derivatives; *therapeutic use]

MeSH check words

Humans